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- (54) Non-A, non-B hepatitis virus particles.
- Disclosed are an isolated non-A, non-B hepatitis virus particle comprising at least one antigen selected from the group consisting of a core antigen, a matrix antigen and an envelope antigen of the non-A, non-B hepatitis virus and a method for efficiently producing the same by genetic engineering. The non-A, non-B hepatitis virus particle can advantageously be used not only for the production of an NANBV hepatitis vaccine exhibiting an extremely high immunogenicity and a diagnostic agent which is extremely high in the antibody detection ratio and in the degree of accuracy of the detection, but is also useful for researches on liver diseases, such as liver cancer.

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# Background of the Invention

#### Field of the Invention

The present invention relates to non-A, non-B hepatitis virus particles and a method for producing the same. More particularly, the present invention is concerned with non-A, non-B hepatitis virus particles which are obtained by expressing the nucleotide sequence of a region selected from the entire region of non-A, non-B hepatitis virus genome, the entire region of the ORF thereof and a region of the ORF which is obtained by cutting off NS4 and/or NS5 from the ORF, and also concerned with an effective method for producing the same. The non-A, non-B hepatitis virus particles of the present invention are useful for providing a vaccine for non-A, non-B hepatitis, a diagnostic reagent for non-A, non-B hepatitis and an agent for screening blood for transfusion for preventing post-transfusion hepatitis each of which comprises the non-A, non-B hepatitis virus particles as an active ingredient, and for providing a polyclonal or monoclonal antibody which is prepared by using the non-A, non-B hepatitis virus particles. Thus, the non-A, non-B hepatitis virus particles of the present invention are useful for producing a vaccine, an immunoglobulin, an immunological diagnostic reagent, an agent for use in affinity column chromatography for removing non-A, non-B hepatitis virus from blood for transfusion.

#### Discussion of Related Art

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Definition of non-A, non-B hepatitis virus:

The viral hepatitis is a liver disease caused by the infection of a hepatitis virus. Heretofore, hepatitis A virus, hepatitis B virus and hepatitis D (deita) virus have been isolated and identified. The hepatitis D virus (deltahepatitis virus) is a deficient virus which cannot multiply by itself and requires for its multiplication the co-presence of hepatitis B virus as a helper virus. Therefore, the hepatitis D virus is present only in a patient having hepatitis B. In 1974, it was reported that there were many patients having hepatitis caused by a factor other than the infection with either hepatitis A virus or hepatitis B virus. Such a hepatitis was named "non-A, non-B hepatitis", and researches on the non-A, non-B hepatitis virus have been made extensively and intensively throughout the world. Heretofore, it has been found that a plurality of types of non-A, non-B hepatitis viruses exist. Results of the researches up to now show that the non-A, non-B hepatitis virus is classified into two types according to the infection route: an epidemic hepatitis virus, namely an enterically-transmitted non-A, non-B hepatitis virus which is spread through water and food; and a blood transmitted non-A, non-B hepatitis virus which spreads over the areas of Africa, India and Southeast Asia has been virologically identified, but the blood-transmitted non-A, non-B hepatitis virus has not yet been identified.

Hereinbelow, the blood-transmitted non-A, non-B hepatitis is often referred to simply as "NANB hepatitis", and the blood-transmitted non-A, non-B hepatitis virus is often referred to simply as "NANBV". Current situation of the studies on NANB hepatitis and problems:

With respect to the epidemiology, clinical examination, diagnosis, treatment and prevention of the NANB hepatitis, virological studies have been made in the world by the comparison of NANBV with the other hepatitis viruses, based on the knowledge of diagnostics, histopathology, immunology, molecular biology and the like ["Japan Medical Journal", No. 3320, pp.3-10, 1987; "Igaku-no Ayumi (Progress of medicine)", 151(13), pp.735-923, 1989; "Kan Tan Sui (Liver, Gallbladder, Pancreas)", 21(1), pp.5-113, 1990; "Jikken Igaku (Experimental Medicine)", 8(3), pp.201-233, 1990]. With respect to the NANB hepatitis, the following findings have been reported

- (1) Epidemiology: In Japan, according to the estimation by the Ministry of Health and Welfare, about 60 % of chronic hepatitis patients (namely about 720 thousand patients), about 40 % of hepatocirrhosis patients (namely about 100 thousand patients) and about 40 % of liver cancer patients (namely about 7 thousand patients) are patients having NANB hepatitis. Further, the mortality attributed to the above-mentioned NANB hepatitis reaches 16 thousand per year. In U.S.A., the number of post-transfusion hepatitis patients reaches 150 to 300 thousand per year and 90 % of the post-transfusion hepatitis patients are patients having NANB hepatitis. Further, it is considered that 1 to 6 % of the blood donors are NANBV carriers. Further, it is estimated that in the other countries also, the incidence of NANB hepatitis and the ratio of the NANBV carrier are equal to or higher than those in U.S.A. and Japan. Therefore, prevention, early diagnosis and early treatment of the NANB hepatitis are of global importance.
- (2) Virology: The NANBV heretofore reported comprises an envelope and assumes a viral particle having a spherical shape of about 50 nm in diameter. The taxonomic observations suggest that the known NANBV is a virus similar to a togavirus or a flavivirus, or a virus of new type different from the togavirus or flavivirus. Further, the results of pathological observations of the cytoplasm of hepatocytes of a plurality of chimpanzees

injected with serum of a patient having NANBV hepatitis show that the formation of a tubular structure occurs in the cytoplasm of a hepatocyte of some of the chimpanzees, but does not occur in the cytoplasm of a hepatocyte of the other chimpanzees, and that an intranuclear particle is formed in the cytoplasm of a hepatocyte of some of the chimpanzees. These results and the results of the epidemiological observations, tests on the presence or absence of the chloroform sensitivity and immunological diagnosis suggest that a plurality of types of NANBVs exist (see, for example, "Science", 205, 197-200, 1979, "Journal of Infectious Disease", 148, 254-265, 1983, and "Biseibutsu" (Microorganism), 5(5) 463-475, 1989). The amount of the NANBV present in the blood of a patient having NANB hepatitis is extremely small as compared to either the amount of a hepatitis A virus present in the feces of a patient having hepatitis A or the amount of a hepatitis B virus present in the blood of a patient having hepatitis B. For example, the amount of hepatitis B virus in the blood of the patient is 108 to 10° per ml in terms of Chimpanzee Infectious dose (CID), whereas the amount of NANBV in the blood of the patient is only 104 to 105 per ml in terms of CID (Bradley, D.W.: Research perspectives in post-transfusion non-A. non-B hepatitis, in "Infection, immunity and Blood Transfusion", edited by Dodd, R.Y. & Barker, L.F., published by Alan R. Liss, Inc., New York (1985) pp.81-97). Further, it is known that except for human, there are no animals except chimpanzee that are sensitive to NANBV and that in the cytoplasm of the hepatocyte, a typical tubular structure is occasionally formed by NANBV infection. Since only chimpanzee can be used as an animal for experiment of the NANBV infection, a large number of chimpanzees are required to be used for the study of NANBV. However, the chimpanzee is not easily available and expensive. Therefore, the study of NANBV by, for example, experimental infection by NANBV, identification of NANBV and search for a useful marker for NANBV, is necessarily restricted and delayed. In order to solve these problems, various attempts have been made for the study of NANBV. For example, in an attempt, an NANBV genomic cDNA [referred to as "hepatitis C virus (HCV)"] was cloned from blood plasma of chimpanzees suffering from NANB hepatitis (Science, 244, 359-362, 1989), and it was confirmed that the antigen (referred to as "C-100") obtained by expressing the cDNA exhibited an antigen-antibody reaction with the antibody in the blood of an NANB hepatitis patient (Science, 244, 362-364, 1989). Further, in another attempt, a chimpanzee was not used and an NANBV genomic cDNA was cloned from the blood plasma of NANB hepatitis patients, and it was confirmed that the antigen obtained by expressing the cDNA exhibited an antigen-antibody reaction with the antibody in the serum of an NANB hepatitis patient (Gastroenterologia Japonica, 24, 540-544 and 545-548, 1989). Furthermore, with respect to the cloning of an NANBV genomic cDNA and the determination of the nucleotide sequence thereof and the corresponding amino acid sequence, clones provided by the following institutions are known: Mitsubishi Kasel Corp., Japan (European Patent Application Publication No. 293274), Chiron Corporation, U.S.A. (European Patent Application Publication Nos. 318216, 388232 and 398748), the Research Foundation for Microbial Diseases of Osaka University, Japan (European Patent Application Publication No. 363025 and Journal of Virology, 65, 1105-1113, 1991), Sanwa Kagaku Kenkyusho Co., Ltd., Japan (Japanese Patent Application Laid-Open Specification No. 1-186990), National Cancer Center Research Institute, Japan [Proceedings of the National Academy of Sciences (U.S.A.), 87, 9524-9528, 1990], Jichi Medical School (Japanese Journal of Experimental Medicine, 60, 167-177, 1990), National Institute of Health, Japan [Nucleic Acid Research, 17(24). 10367-10372, 1989; the same literature, 18(15), 4626, 1990; Gene, 91, 287-291, 1990; and Journal of General Virology, 71, 3027-3033, 1990), and the like. Moreover, concerning the structure of NANBV gene, it has been reported: that the total length of NANBV genome is about 10 kb; that the genome is comprised of a non-coding region at the 5'-end, an open reading frame (ORF) region and a non-coding region at the 3'-end; that in the ORF region, genes which code for a virus core antigen (protein) (C antigen), a matrix antigen (protein) (M antigen), an envelope antigen (protein) (E antigen), and six types of non-structural proteins (NS proteins) are disposed in this order from the 5'-end to the 3'-end; and that the NS protein gene is comprised of NS1, NS2, NS3, NS4a, NS4b and NS5 which are disposed in this order from the 5'-end to 3'-end. With respect to the functions of these antigens (proteins) it is believed that C antigen is responsible for the protection of the gene, E antigen is responsible for infection, M antigen is responsible for the maintenance of the structure of E antigen, NS1 serves as a complement fixing antigen, NS3 serves as a protease, NS5 serves as a polymerase and the noncoding region is responsible for the maintenance of the structure of the genome and for the replication of the genome. The functions of NS2 and NS4 have not yet been known.

(3) Clinical observations: Hepatitis is generally classified either into epidemic hepatitis and sporadic hepatitis according to the number and frequency of the occurrences of hepatitis, or into acute hepatitis, fulminant hepatitis, subacute hepatitis, persistent hepatitis and chronic hepatitis according to the severeness and stage of the hepatitis patients. The latent period of the NANB hepatitis is 2 to 26 weeks. The symptom of NANB hepatitis in the early stage is mild as compared to that of hepatitis B. For example, a patient having NANB hepatitis only becomes feverish and complains of languor. Further, 70 % of the patients have anicteric symptom. Therefore, the NANB hepatitis is frequently overlooked. However, the NANB hepatitis is very dangerous because the NANB hepatitis is likely to become chronic and, then, to progress to liver cirrhosis. Illustratively stated, 40

to 50 % of the patients having NANB hepatitis whose serum exhibits an increased aminotransferase activity develop chronic hepatitis. 10 to 20 % of the cases of chronic hepatitis suffer from liver cirrhosis. Further, 0.5 to 1 % of blood recipients per year becomes liver cirrhosis patients without subjective symptoms. More seriously, the liver cirrhosis may further progress to liver cancer or hepatoma. Therefore, for preventing biohazard caused by blood transfusion and bleeding, eradication of the NANB hepatitis is a matter of global importance from the viewpoint of public health.

(4) Diagnosis: As mentioned above, the NANBV (blood-transmitted type) has not yet been identified and a viral marker, such as an NANBV antigen, which is useful for the diagnosis of NANB hepatitis has not been known. Therefore, diagnosis of NANB hepatitis has been conducted by examining the titer of the antibody in serum of a patient, which is specific for each of the known pathogenic viruses, such as hepatitis A virus, hepatitis B virus, cytomegalovirus, EB virus, varicella virus and herpes simplex virus, and diagnosing the patient whose serum is negative with respect to the antibody specific for any of the above-mentioned viruses, as having NANB hepatitis, or by performing a histopathological examination through a biopsy of the liver ("Disease of the Liver and biliary system", 8th edition, S. Shenlock, pp. 326-333, 1989, Blackwell Scientific Publications). At the same time, another diagnosis method has also been used. For example, there have been used a method in which the activity of an enzyme in serum, such as GPT [glutamic-pyruvic transaminase, also known as "ALT" (alanine aminotransaminase)], GOT [glutamic-oxalo-acetic transaminase, also known as "AST" (aspartate aminotransferase)], and guanine deaminase (also known as "guanase") is determined ("Kan Tan Sui (Liver, Galibladder, Pancreas)", Vol. 14, pp. 519-522, 1987). With respect to the GPT or GOT in serum mentioned above, a standard for the diagnosis of NANB hepatitis in which lasting and abnormally high activities of GPT and GOT are utilized as a criterion for the diagnosis of NANB hepatitis, is employed in Japan ("Journal of Blood Transfusion Society in Japan", 31(4), 316-320, 1985; and "Nippon Rinsho", 46, 2635-2638, 1988). Regarding the immunological diagnosis, in the present situation in which the isolation and identification of NANBV are difficult as described above, an antigen-antibody reaction between an antigen obtained by expression of NANBV cDNA clone (which has been isolated using the techniques of genetic engineering and the knowledge of immunology) and the serum of an NANB hepatitis patient is used as a criterion. Examples of known antigens include an expression product of an NANBV cDNA prepared from the plasma of an NANB hepatitis patient (European Patent Application Publication No. 363025), an expression product of "HCV" cDNA prepared from the plasma of a chimpanzee having the symptoms of NANB hepatitis (European Patent Application Publication No. 318216 and Japanese Patent Application Laid-Open Specification No. 2-500880), an expression product of an NANBV cDNA derived from the liver of an NANBV-infected chimpanzee (European Patent Application Publication No. 293274, Japanese Patent Publication Specification No. 64-2576 and Japanese Patent Application Laid-Open Specification No. 1-124387). As a method for determining the antigen-antibody reaction, RIA (radioimmunoassay) and EIA (enzyme immunoassay) are generally used. However, these expression products are different in antigenicity. The antigen which is an expression product of HCV cDNA (that is, the C-100 antigen mentioned) can be some criterion or yardstick for the diagnosis of chronic hepatitis caused by the HCV infection. However, since the region in which the antigen (C-100) exhibits its antigenicity is limited and the detection ratio of the antibody is as disadvantageously low as about 70 % ["Biseibutsu (Microorganism)", 5, 463-475, 1989; "Kan Tan Sui (Liver, Gallbladder, Pancreas)", 20, 47-51, 1990; and "Igaku-no Ayumi (Progress of Medicine)", 151, 871, 1989], this antigen is unsatisfactory from the viewpoint of accurate diagnosis of NANB hepatitis and NANBV infection and from the viewpoint of accurate determination of the progress of a patient suffering from chronic hepatitis and acute hepatitis for treatment thereof. Therefore, it has been desired to obtain a reliable method for the diagnosis and prognosis of the NANB hepatitis.

(5) Therapy and Prevention: Recently, the usefulness of  $\alpha$ - and  $\beta$ -interferons in the treatment of chronic NANB hepatitis have been reported (\*Kan Tan Sui (Liver, Gallbladder, Panceras)\* vol. 20, pp. 59-64, 1990; "Igaku-no Ayumi (Progress of Medicine)\*, vol. 151, pp. 871-876, 1989). However, a suitable dose of  $\alpha$ - and  $\beta$ -interferons and a suitable period for administration thereof have not yet been established.

On the other hand, for prevention of NANB hepatitis, various vaccines are used in which the above-mentioned conventional expression products of NANBV cDNAs (European Patent Application Publication No. 363025) or HCV cDNAs (European Patent Application Publication No. 318216) are used as an antigen. However, as is apparent from the fact that the NANBV itself has not yet been isolated and identified before completion of the present invention, it has been impossible to specify an antigen useful for NANBV vaccines from the above-mentioned expression products each having a variety of antigenic determinants (epitopes) and determine the effectiveness and safety of such a specific antigen so that the antigen can be clinically used. Accordingly, there is no NANBV vaccine which can be advantageously put into practical use.

(6) Production of NANBV particles and significance thereof: although various NANBV cDNA clones have been known as described in item (2) above, no report has been made such that an NANBV particle has been successfully produced by using the known clones. This fact means that it is extremely difficult to construct a

cDNA of about 10 kb which covers the entire region of NANBV genome, the cDNA being necessary for the production of an NANBV particle. That is, by using the prior art technique for the selection of materials to be used for the extraction of NANBV genomic RNA and the prior art technique for the extraction and the purification of the RNA, it is only possible to isolate a short RNA fragment of at most a few hundreds nucleotides and a cDNA clone thereof. When it is intended to construct a cDNA of the entire region of NANBV genome by using such a short-length cDNA fragment, it is necessary to select more than several tens of different types of cDNA fragments in a combination such that the ORFs of the cDNA fragments can form the entire region of NANBV genome, and ligate them in sequence accurately without any mistake. Needless to say, the operation for ligating cDNA fragments in sequence while satisfying such strict requirements is extremely cumbersome and difficult. It should be noted that the probability of the occurrence of a fatal mistake in the ligating operation for cDNA fragments is increased in proportion to the increase in the number of ligations. Therefore, in order to attain the accurate construction of the cDNA of the entire region of NANBV genome, it is necessary to reduce the number of ligation steps by using cDNA fragments which are as long as possible. It should further be noted that the realization of the reduction of the number of the ligations needs a high level of academic knowledge and experience and extraordinary skills with respect to the preparation by extraction of a long-length NANBV genomic RNA fragment of about 2 kb to about 5 kb and with respect to the cloning of a cDNA thereof. On the other hand, as described in item (4), since the antigen used in the commercially available diagnostic reagents for NANB hepatitis is an expression product of a part of NANBV genomic cDNA fragment and, therefore, is narrow in the antigen spectrum, the antigen reacts mainly with the serum of a chronic hepatitis patient and exhibits an antibody detection ratio as low as about unsatisfactory 70 %. Therefore, a diagnostic reagent is in a great demand which exhibits excellent specificity in the antigen-antibody reaction with the serum of not only a chronic NANB hepatitis patient but also an acute NANBV hepatitis patient and is high in the antibody detection ratio. To meet the demand, it has been earnestly desired to develop a diagnostic reagent employing, as an antigen, for example, an NANBV particles having a broad antigen spectrum and which exhibits a high detection ratio for antibody. Further, the production of NANBV particles is considered to contribute to solving the problem of item (5) so that a practically employable NANB hepatitis vaccine can be realized. From the foregoing it is apparent that the construction of a cDNA of the entire region of NANBV genome and the attainment of the mass production of NANBV particles by expressing the cDNA have been earnestly desired as a matter of global interest.

#### Summary Of The Invention

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The present inventors have made extensive and Intensive studies with a view toward solving the abovementioned problems of the prior art by developing novel isolated NANBV particles. As a result, the present inventors have succeeded in constructing an ORF (open reading frame) region from the C antigen gene through the NS3 gene of the NANBV genomic cDNA, an entire ORF region of the NANBV genomic cDNA which is longer than the above-mentioned ORF region, and the entire region of NANBV genome comprised of the above entire ORF region and, ligated at its 5'-end and 3'-end, non-coding regions for 5'-end and 3'-end, by skillfully ligating not more than ten different NANBV cDNA clones each comprising at least 1000 nucleotides so that a desired ORF having or not having non-coding regions at 3'- and 5'ends is constructed. Moreover, the present inventors have surprisingly succeeded in the mass production of NANBV particles by introducing or inserting each of the above-mentioned regions of the NANBV genome individually into an expression vector and expressing the regions. In the present invention, the terminology "non-A, non-B hepatitis virus particle" means an expression product of the above-mentioned regions of the NANBV genome and comprises at least one antigen selected from the group consisting of a core antigen, a matrix antigen and an envelope antigen of the non-A, non-B hepatitis virus. Examples of non-A, non-B hepatitis virus (NANBV) particles include those of the following structures: a complete NANBV particle which is an NANBV antigen assembly comprised mainly of C (core) antigen, M (matrix) antigen and E (envelope) antigen and which has a nucleic acid in the virus particle; an incomplete NANBV particle which is an NANBV antigen assembly comprised mainly of C antigen, M antigen and E antigen but which has no nucleic acid in the virus particle; an NANBV core which is an NANBV antigen assembly comprised mainly of C antigen and which has a nucleic acid in the core; an incomplete NANBV core which is an NANBV antigen assembly comprised mainly of C antigen but which has no nucleic acid in the core; and an NANBV surface antigen assembly comprised mainly of E antigen. This success is attributed to a unique technique of the present inventors such that in order to obtain an authentic NANBV genome, NANBV RNAs are extracted directly from NANBV particles contained in whole blood of a patient having NANB hepatitis or a resected liver of a patient having NANB hepatitis and liver cancer in combination, without multiplying the NANBV in a chimpanzee having unknown factors which are considered to have rendered difficult the isolation of NANBV, although the amount of NANBV in the blood or resected liver is extremely small, that is as small as about 1/10,000 that of a hepatitis A virus or a hepatitis B virus, but with paying minute care in the operating procedure so that the NANBV and

its genome do not undergo cleavage and/or decomposition by the action of body fluids or blood enzymes during the storage of fresh materials for NANBV genome and that a complete NANBV genomic RNA or RNA fragments having a length of about 2 kb to 5 kb are obtained. RNAs thus prepared from fresh human materials are then converted to cDNA by means of a reverse transcriptase to obtain a cDNA library. In order to screen a NANB genomic cDNA of about 1000 to about 5000 nucleotides from the cDNA library, the cDNAs are individually inserted in lambda gt11 phage vectors and then expressed on the phage plaques at high concentration, followed by screening of NANBV genomic cDNAs by repeatedly conducting enzyme immunoassay (EIA) in which both serum from a convalescent patient having acute NANB hepatitis and serum from a patient having chronic NANB hepatitis are used. Thus, safe production of NANBV particles or NANBV antigen assemblies with high purity on a large scale at low cost without biohazard, has for the first time been realized by expressing by recombinant DNA techniques the entire region of NANBV genomic cDNA or the entire ORF region of NANBV genomic cDNA constructed by selecting cDNA clones covering the entire region or the entire ORF region of NANBV genomic cDNA, cutting-off any overlapping portion from the cDNA clones and ligating the cDNA clones in sequence such that the entire region or the entire ORF region of NANBV genomic cDNA is formed. Furthermore, it has been found that the expression product of the present invention has an extremely broad antigen spectrum as compared with a conventional expression product of NANBV genomic cDNA fragments of short length, and exhibits antigen-antibody reaction specifically with the serum of both a chronic patient and an acute patient of NANB hepatitis so that the detection ratio for the antibody is 95 % or more, solving the problem of item (6) above. That is, the expression product of the present invention has been found to make a great contribution to the prevention, diagnosis and treatment of NANB hepatitis by providing a vaccine having an enhanced immunogenicity, a diagnostic reagent exhibiting an improved detection ratio for antibody and an improvement in the preparation of antibodies. Based on the above, the present invention has been completed.

The foregoing features and advantages of the present invention will be apparent from the following detailed description and appended claims taken in connection with the accompanying drawings.

## **Brief Description Of The Drawings**

In the Drawings:

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Fig. 1(1) and Fig. 1(2) are diagrams showing the relationships between the cDNA clones of the NANBV gene to be used in the present invention, shown relative to the entire region of the NANBV genome; Fig. 2(1) through Fig. 2(16) show the nucleotide sequence of the entire region of the NANBV genomic cDNA to be used in the present invention and the amino acid sequence coded for by the nucleotide sequence; Fig. 3 is a diagram showing the hydrophobicity profiles of both of the NANBV of the present invention and the Japanese encephalitis virus (JEV), in which the hydrophobicity index of the NANBV is compared with that of the JEV, and wherein the abscissa indicates the amino acid number, the ordinate indicates the hydrophobicity index, the vacant triangle indicates the glycosylation site, the asterisk indicates the site of amino acid sequence (Gly-Asp-Asp) common to RNA polymerase, and C, M, E and NS represent core antigen, matrix antigen, envelope antigen and non-structural protein, respectively.

Fig. 4 is a diagram showing the steps for the construction of plasmid pMAM-neo10 for expressing the NANBV genomic cDNA in an animal cell.

Fig. 5 is a diagram showing the steps for the construction of plasmid pYHC5 for expressing the NANBV genomic cDNA in yeast.

Fig. 6 is a diagram showing the steps for the construction of plasmid pXX-49, pXX-51 and pXE-39 for the preparation of a recombinant vaccinia virus.

Fig. 7 is a graph showing the sucrose concentration and the antigenic activity of each of the fractions obtained by sucrose density-gradient centrifugation of the supernatant of the culture of recombinant vaccinia virus vXX39.

Fig. 8 is an electron microscopic photomicrograph of NANBV particles produced by culturing cells infected with recombinant vaccinia virus vXX39.

# Detailed Description Of The Invention

Essentially, according to the present invention, there is provided an isolated non-A, non-B hepatitis virus particle comprising at least one antigen selected from the group consisting of a core antigen, a matrix antigen and an envelope antigen of the non-A, non-B hepatitis virus.

In a preferred embodiment of NANBV particle of the present invention, the core antigen, the matrix antigen and the envelope antigen are, respectively, coded for by a nucleotide sequence of the 333rd to 677th nucleotides, a nucleotide sequence of the 678th to 905th nucleotides and a nucleotide sequence of the 906th to

1499th nucleotides of the non-A, non-B hepatitis virus entire nucleotide sequence from the 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2(16) hereof.

In another aspect of the present invention, there is provided a non-A, non-B hepatitis virus particle" which has a ribonucleic acid corresponding to at least part of the nucleotide sequence shown in Fig. 2(1) through Fig. 2(16) hereof.

In the present invention, unless otherwise specified, the left end and right end of-the sequence of deoxyribonucleotides are the 5'-end and 3'-end, respectively. Further, unless otherwise specified, the left end and right end of the amino acid sequences of peptides are the N-terminus and C-terminus, respectively.

The isolated NANBV particle of the present invention can be prepared and identified in accordance with the following steps (I) to (IX).

Step (I): Selection and collection of a material for extracting an NANBV genomic RNA.

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As a material for extracting the NANBV RNA, there may be used, for example, blood, lymph, ascites and hepatocyte of an NANBV carrier, or of a human or a chimpanzee suffering from NANB hepatitis, and hepatocyte of a patient suffering from NANB hepatitis and liver cancer or hepatoma in combination. Since the materials derived from a chimpanzee may contain NANBV in a relatively small amount as compared to the materials derived from a human and a chimpanzee has unknown factors which are considered to have rendered difficult the isolation of NANBV, the use of the materials derived from a human is preferred. Of blood, lymph, ascites and hepatocytes from a human, blood can most easily be obtained in a large amount. For example, blood which is not acceptable for use as blood for transfusion is available in a large, amount from, e.g., a blood bank. Such blood can advantageously be used as a material for extracting an NANBV RNA. When blood is used as a material, blood is separated into plasma and erythrocytes. The thus obtained plasma is examined to determine whether or not the plasma is negative to the surface antigen of hepatitis B virus (WHO expert committee on viral hepatitis: Advances in viral hepatitis, WHO Technical Report Series, 602, 28-33, 1977) and negative to a genomic DNA of hepatitis B virus (Brechot, C., Hadchouel, M., Scotto, J., Degos, F., Charnay, P., Trepo, C., Tiollals, P.: Detection of hepatitis B virus DNA in liver and serum: a direct appraisal of the chronic carrier state. Lancet 2: 765-768, 1981). Further, the plasma is examined with respect to the activities of enzymes, such as GPT (Wroblewski, F. & LaDue, J. S.: Serum glutamic-pyruvic transaminase in cardiac and hepatic disease, Proc. Soc. Exp. Biol. Med., 91, 569, 1956), GOT, guanase and the like, which are employed as the criterion for the diagnosis of NANB hepatitis. The above-mentioned procedures of the separation of blood into plasma and erythrocytes and the examination of the plasma are conducted with respect to blood of different lots. The plasma which is negative to both surface antigen and genomic cDNA of hepatitis B virus and exhibits extremely high activities of the above-mentioned enzymes, for example, a GPT activity of 35 IU/ml or more, is pooled.

The number of the NANB hepatitis virus particles in blood is extremely small as compared to that of the hepatitis B virus particles as mentioned hereinbefore. From the results of the infection experiment, the number of the NANB hepatitis virus particles in blood is estimated to be about 1/10,000 of the number of the hepatitis B virus particles (Bradley, D.W., (1985): Research perspectives in post-transfusion non-A, non-B hepatitis, in "Infection, Immunity and Blood Transfusion", edited by Dodd, R.Y. & Barker, L.F., published by Alan R. Liss, Inc., New York, pp. 81-97). Therefore, for the extraction of the RNA, it is preferred to use blood in a large amount, for example, in an amount as large as about 3 to 10 liters. Fresh whole blood to be used as a material for extracting an NANB RNA from NANBV particles is stored at 1 to 5 °C in order to prevent NANBV and its gene from being denatured and to prevent its gene from being cleaved or decomposed by the action of an enzyme. It is also desirable to complete the preparation of NANBV RNAs by Step (II) within 48 to 72 hours from the collection of the fresh whole blood. When a hepatocyte is used as a material, about 1 to 3 g of a non-cancerous or a cancerous portion of a liver tissue resected from a patient having hepatoma or liver cancer which is a complication of a chronic NANB hepatitis may advantageously be used. Hepatocyte to be used as a material is stored in a frozen state at -70 °C.

Step (II): Preparation of the NANBV RNA

From the material obtained in Step (I), the RNA may be extracted and purified by conventional methods. For example, when fresh whole blood is used as the material, about 3 to 10 liters of fresh whole blood is subjected to low-speed centrifugation to collect a plasma fraction as a supernatant. The virus fraction is obtained from the plasma through purification for use in the subsequent procedure for the extraction and purification of the RNA.

On the other hand, when hepatocyte is used as a material for extracting the NANBV RNA, about 5 to 30-fold volume of a diluent containing ribonuclease inhibitor is added to the liver tissue. Then, according to the con-

ventional method using a homogenizer and the like, the liver tissue is crushed or disrupted to obtain a homogenate of hepatocyte. As a diluent, 10 to 150 mM of a conventional buffer may be used. Then, the homogenate is subjected to low-speed centrifugation to collect a supernatant. The collected supernatant is used as an original solution for the extraction and purification of the NANBV RNA. The extraction and purification of the NANBV RNA may be conducted by the conventional method, for example, an extraction method in which a mixture of a ribonuclease inhibitor, such as heparin, diethyl pyrocarbonate, and guanidine thiocyanate, with a surfactant, a chelating agent, or a reducing agent capable of enhancing the denaturation of a protein, is used; a method in which fractionation is conducted by density gradient centrifugation using sucrose, cesium chloride, cesium trichloroacetate, Ficoli (Pharmacia Fine Chemicals AB, Sweden) or the like as a solute of a gradient; a method in which separation is conducted by affinity column utilizing the 3'-terminal poly A chain which an mRNA specifically has; a separation method in which an mRNA-bonded polysome is obtained by the immunoprecipitation using an antibody specific for a protein synthesized on the polysome; a phenol extraction method based on a principle of two-phase separation; a precipitation method by the use of a polyethylene glycol, a dextran sulfate, an alcohol or the like. The above-mentioned methods may be used individually or in combination. The abovementioned procedure for extracting and purifying the NANBV RNA may preferably be conducted at pH 3 to 10 in order to prevent the irreversible denaturation of the RNA. Thus, NANBV RNAs are obtained.

Step (III): Preparation of a double-stranded cDNA from the NANBV RNA

Using as a template each of the NANBV RNAs abtained in Step (II), a cDNA may be prepared by a standard method. That is, using an oligodeoxythymidine and a random hexanucleotide primer as primers and using a reverse transcriptase, a cDNA complementary to the NANBV RNA is synthesized using the NANBV RNA as a template to obtain a double-strand comprising the cDNA and the NANBV RNA which are complementarily bended to each other. Then, the thus obtained double-strand is reacted with ribonuclease H so that the NANBV RNA is decomposed and removed from the cDNA. Thus, a single-stranded cDNA is obtained. Using the obtained single-stranded cDNA as a template, a double-stranded cDNA is synthesized by means of a DNA polymerase. The double-stranded cDNA synthesis may easily be conducted using a commercially available kit for cDNA synthesis, for example, cDNA Synthesis System Plus® (manufactured and sold by Amersham, England; BRL Inc., U.S.A.), cDNA System Kit® (manufactured and sold by Pharmacia LKB, Sweden), cDNA Synthesis Kit® (manufactured and sold by Boehringer Mannheim GmbH, Germany), and the like. When the quantity of the synthesized cDNA is small, the cDNA can be amplified using a conventional method, such as PCR (polymerase chain reaction) method ("PCR Technology", edited by H.A. Erlich, published by Stockton Press, 1989) using a PCR kit, such as AmpliTaq (manufactured and sold by Perkin Elmer Cetus, U.S.A.). Thus, double-stranded cDNAs are obtained.

Step (IV): Preparation of a cDNA library

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Using the cDNAs prepared in Step (III), a cDNA library is prepared by a customary method. That is, the cDNAs prepared in Step (III) are individually ligated to replicable cloning vectors, to thereby obtain a cDNA library. As a replicable cloning vector, any known or commercially available vectors, such as phage, cosmid, plasmid and animal virus may be used. When a phage or a cosmid is used as a replicable vector, in order to attain high stability and high transforming ability of the vector after each of the cDNA fragments has been individually inserted therein, the in vitro packaging of each of the cDNA-inserted vectors is conducted by a customary method. Thus, the cDNA-inserted vectors are obtained in the form of a recombinant phage particle. The obtained phage particles are used as a cDNA library for cDNA cloning. On the other hand, when a plasmid is used as a replicable vector, the above-mentioned cDNA fragments are individually inserted in the plasmid vectors and the resultant cDNA-inserted vectors are then individually introduced into sensitive host cells, such as cells of Escherichia coli, Bacilus, subtilis, yeast or the like, according to a customary method. The thus obtained transformants are used as a cDNA library for cDNA cloning. Further, when the animal virus gene is used as a replicable vector, the above-mentioned cDNA fragments are individually inserted in the virus vectors and the resultant recombinant viruses are then individually infected into sensitive animal cells according to a standard method and multiplied in the cells. In the case of the recombinant virus, the obtained recombinant viruses as such are used as a cDNA library.

The preparation of the cDNA library may easily be conducted using a commercially available kit, for example, a cDNA cloning system lambda gt10 and lambda gt11 (manufactured and sold by Amersham, England; BRL Inc., U.S.A.; and Stratagene Inc., U.S.A.), an in vitro packaging system (manufactured and sold by Amersham, England; BRL Inc., U.S.A.; and Stratagene Inc., U.S.A.) and the like.

Step (V): Cloning of a cDNA clone containing an NANBV gene from the cDNA library

In this step, a cDNA clone containing an NANBV gene is obtained. When the cDNA library is comprised of transformants, the transformants are cultured on a standard agar medium to form colonies. On the other hand, when the cDNA library is comprised of recombinant phage particles or recombinant viruses, these phage particles or recombinant viruses are used to infect known sensitive host cells, such as Escherichia coli, Bacillus subtilis, yeast, animal cell culture and the like, and cultured to form plaques, or to multiply the infected cells. The above-obtained transformant colonies, plaques or infected cells are subjected to immunoassay by at least one of the standard methods individually using serum from a convalescent patient having acute NANB hepatitis, serum from a patient having chronic NANB hepatitis, and serum from chimpanzee infected with an NANBV irrespective of whether or not the NANBV is of the type which causes a tubular structure to be formed in the cytoplasm of the hepatocyte of the chimpanzee, so that colonies, plaques or infected cells which have produced an NANBY antigen specifically reacted with at least one of the above-mentioned sera are selected and isolated. For the strict selection of the colonies, plaques and infected cells, it is preferred that the above procedure be repeated. From each of the thus selected and isolated colonies, plaques or the infected cells, a cDNA clone containing an NANBV gene is isolated according to a standard method described in T. Manlatis et al., Molecular Cloning, A Laboratory Manual, published by Cold Spring Harbor Laboratory, U.S.A., pp. 309-433 (1982). The immunoassay may be conducted by, for example, an enzyme-labeled antibody technique in which an antibody labeled with an enzyme, such as peroxidase and alkaline phosphatase is used; and a fluorescent antibody technique in which an antibody labeled with fluorescein isothiocyanate, europium or the like is used. It is preferred that the immunoassay by the above-mentioned technique be conducted by an indirect method because with the indirect method, high sensitivity immunoassay can be attained even by the use of an extremely small amount of serum from a patient. As a primary antibody to be used in the indirect method, serum from a patient having NANB hepatitis or serum from a chimpanzee having NANB hepatitis may preferably be employed because these sera contain an antibody specific for an NANBV antigen in relatively large amount. As a secondary antibody to be used in the indirect method, a commercially available anti-human Ig (immunoglobulin) antibody labeled with an enzyme, a fluorescent substance or the like may be used.

A specimen to be subjected to immunoassay may be prepared according to a conventional method, for example, a blotting method in which nucleic acids and proteins of the colonies, plaques and infected cells are adsorbed on a filter membrane, a method in which a microplate or a slide glass for microscopy is used, or the like. When the blotting method is used in combination with an indirect, enzyme-labeled antibody technique, the selection of the intended colonies, plaques or infected cells from an extremely large number of the original colonies, original plaques or original infected cells can be conducted easily and promptly. In this case, blotting is conducted by contacting a commercially available filter made of nitrocellulose, cellulose acetate, nylon or the like, with the colonies, plaques or infected cells.

The above-obtained cDNA clone is a part of the NANBV gene. Therefore, in order to obtain cDNA clones covering the entire region of the NANBV gene, it is requisite to extend the cNDA clone by a method in which cDNA fragments adjacent to the cDNA clone are isolated by using 3'- and 5'- terminals of the cDNA clone as a probe. In this case, the technique which is known as "gene walking" (also known as "genomic walking" or "chromosome walking") may be employed ("DNA cloning volume III", edited by D.M. Glover, pp.37-39, IRL Press, 1987; "Molecular Cloning - a laboratory manual" 2nd edit., T. Maniatis et al, 3.21 - 3.23, 1989). By the repetition of the doning procedure and the gene walking, the entire region of the NANBV gene can be obtained in the form of cDNA clones.

Further, the nucleotide sequence of each of the obtained cDNA clones is determined. The determination of the nucleotide sequence of the cDNA clone may be conducted according to a conventional method, for example, the Maxam-Gilbert method, the dideoxy chain termination method (Analytical Biochemistry, 152, 232-238, 1986); or the like.

Based on the determined nucleotide sequence, the amino acid sequence can be deduced. The sequencing of the amino acids is conducted from the location of the initiation codon (ATG on the cDNA or AUG on the mRNA). Important portions of the amino acid sequence, for example, a hydrophilic portion, which is considered to constitute an epitope, can be identified by synthesizing a peptide corresponding to each hydrophilic portion and purifying the synthesized polypeptide by HPLC (high performance liquid chromatography), followed by subjecting the purified peptide to EIA (enzyme immunoassay) or RIA (radioimmunoassay).

The cDNA clones are preferably classified into groups according to the respective properties of the NANBV antigens coded for by the cDNA clones in order to distinguish clones from one another. In this connection, the location of each cDNA clone on the restriction map of the NANBV gene can be used as a yardstick for the classification (see Fig. 1(1) and Fig. 1(2)]. Further, it has been found that some of NANBVs have the ability to cause a tubular structure to be formed in the cytoplasm of a hepatocyte of a chimpanzee, and some of NANBV do

not have such ability (Science, 205, pp. 197-200, 1979). Therefore, the cDNA clones may be identified and classified by examining the serological reactivity of each cDNA clone with serum from a chimpanzee infected with an NANBV of the type which causes a tubular structure to be formed in the cytoplasm of the hepatocyte of the chimpanzee and with serum from a chimpanzee infected which an NANBV of the type which does not cause a tubular structure to be formed in the cytoplasm of the hepatocyte of the chimpanzee. The examination of this serological reactivity may be conducted by immunoassay mentioned above.

In the present invention, as shown in Figs. 1(1) and 1(2), the cDNA clones of the NANBV gene to be used in the present invention are identified with prefix "BK".

Fig. 1(1) is a diagram showing the relationships between the cDNA clones of the NANBV gene to be used in the present invention, shown relative to the entire region of the NANBV gene, and Fig. 1(2) is a diagram showing the relationships between the cDNA clones obtained by gene walking, shown relative to the entire region of the NANBV gene.

These BK NANBV cDNA clones include, for example, Escherichia coli BK 108 (deposited at Fermentation Research Institute, Japan under the accession number FERM BP-2971), Escherichia coli BK 129 (deposited at Fermentation Research Institute, Japan under the accession number FERM BP-2972), Escherichia coli BK 138 (deposited at Fermentation Research Institute, Japan under the accession number FERM BP-2973), Escherichia coli BK 153 (deposited at Fermentation Research Institute, Japan under the accession number FERM 2974), Escherichia coli BK 157 (deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3243), <u>escherichia coli</u> BK 166 (deposited at Fermentation Research Institute, Japan under the accession number FERM BP-2975), and Escherichia coli BK 172 (deposited at Fermentation Research Institute, Japan under accession number FERM BP-2976). These seven BK NANBV cDNA clones are considered to cover at least the entire region of the open reading frame of the NANBV gene and probably the entire region of the NANBV gene (see Fig. 1(1) and Fig. 1(2) hereof). Further, in addition to the above-mentioned cDNA clones, the following five clones are deposited at Fermentation Research Institute, Japan as representative ones of the BK NANBV cDNA clones: Escherichia coli BK 102 (deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3384), Escherichia coli BK 106 (deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3385), Escherichia coli BK 112 (deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3386), Escherichia coli BK 146 (deposited at Fermentation Research Institute, Japan under the accession number FERM-3387), and Escherichia coli BK 147 (deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3388).

The nucleotide sequence of the entire region of the NANBV gene which is covered by the above-mentioned BK NANBV cDNA clones and the amino acid sequence which is coded for by this nucleotide sequence are shown in Fig. 2(1) through Fig. 2(16). Based on the entire NANBV nucleotide sequence and the entire NANBV amino acid sequence shown in Fig. 2(1) through Fig. 2(16), various studies and observations can be made with respect to the homology of the nucleotide sequence and amino acid sequence of the NANBV gene to those of other virus genes, the hydrophobicity index shown in Fig. 3 (hydrophobicity/hydrophilicity profile), the structure of the NANBV gene, the regions of epitopes (antigenic determinants) and the like.

With respect to the homology, studies can be made by comparison of the nucleotide sequence and amino acid sequence of the NANBV gene with those of various viruses whose genes are well known (Japanese Patent Application Laid-Open Specification No. 62-286930 and "Virology", Vol. 161, pp. 497-510, 1987) and those of other viruses, such as bovine virus diarrhea-mucosal disease virus ("Virology", Vol. 165, pp. 497-510, 1988), swine cholera virus ("Virology", Vol. 171, pp. 555-567, 1989), tobacco vein mottling virus ("Nucleic Acid Research, Vol. 165, pp. 5417-5430, 1986), etc.

With respect to the analysis of the hydrophobicity index, studies can be made by techniques using, for example, a genetic information processing software, SDC-Genetyx (manufactured and sold by SDC Software Co., Ltd., Japan), Doolittle's program (Journal of Molecular Biology, Vol. 157, pp. 105-132, 1982) and the like.

The regions of the NANBV gene coding for the various antigens (proteins) of the NANB virus particle, that is, three structural proteins, namely, virus core antigen (protein) (C antigen), matrix antigen (protein) (M antigen) and envelope antigen (protein) (E antigen) and six non-structural proteins (NS proteins) can be determined by comparing the peptides coded for by the genes with known flavivirus with respect to the hydrophobicity index and comparing the amino acid sequences of the peptides with the peptide linking sites which are acted on by the signal peptidase derived from the' host cell (Journal of Molecular Biology, 167, 391-409, 1983) and the serine protease derived from the known flavivirus (virology, 171, 637-639, 1989). With respect to the NANBV particle of the present invention, the antigens (proteins) are, respectively, coded for by the following nucleotide sequences shown in Fig. 2(1) through Fig. 2(16):

C antigen: from the 333rd to 677th nucleotides M antigen: from the 678th to 905th nucleotides

E antigen: from the 906th to 1499th nucleotides
NS1 protein: from the 1500th to 2519th nucleotides
NS2 protein: from the 2520th to 3350th nucleotides
NS3 protein: from the 3351st to 5177th nucleotides
NS4a protein: from the 5178th to 5918th nucleotides
NS5 protein: from the 6372nd to 9362nd nucleotides

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These nucleotide sequences are useful for the diagnosis of NANB hepatitis. The antigens (proteins) which are respectively coded for by these nucleotide sequences are useful as antigens for not only vaccines but also diagnostic reagents for NANB hepatitis. Furthermore, it has been found that since the NANBV particle of the present invention has various types of epitopes, a diagnostic reagent using the NANBV particle or NANBV antigen assembly of the present invention as an antigen has a broad antigen-antibody reaction spectrum and therefore can react to a wide variety of antibodies produced by infection with NANB hepatitis virus, as compared to an antigen containing a single epitope, so that it has high sensitivity in detecting NANB hepatitis, as shown in the Examples described later.

Step (VI): Expression of the entire region of the NANBV genomic cDNA and the ORF thereof and a mass production of an NANBV antigen assembly, an incomplete NANBV particle and an infective, complete NANBV particle.

In order to express the NANBV genomic cDNA cloned in Step (V) and produce an NANBV particle on a commercial scale, part or whole of the cloned cDNA present in the cDNA clone is taken out by cutting from the replicable cloning vector and recombined with a replicable expression vector. Illustratively stated, part or whole of the cDNA of each cDNA clone is taken out by cutting with a restriction enzyme to obtain an cDNA fragment containing an NANBV antigen gene (hereafter referred to as "NANBV DNA fragment"). The NANBV DNA fragments are ligated in sequence so that the entire region of the NANBV gene or the entire region of the ORF thereof is constructed and then inserted in a replicable expression vector. In order to simplify the ligating procedure for the cloned NANBV DNA fragments and prevent the occurrence of a mistake in the ligation, not greater than ten different, preferably not greater than five different NANBV DNA fragments are used for the construction of the entire region of NANBV genomic cDNA or the entire region of the ORF thereof. To realize this, NANBV DNA fragments covering the entire region of the NANBV gene or the entire region of the ORF thereof and each having a length of at least 1000 nucleotides, preferably, 1500 nucleotides, are strictly selected and any overlapping between the fragments is deleted and then the NANBV DNA fragments are ligated in sequence to thereby construct the entire region of the NANBV gene or the entire region of the ORF thereof. That is, it is necessary to provide not more than ten different cDNA clones each comprising at least 1000 nucleotides and prepared from a NANBV genomic RNA fragment of at least 1000 nucleotides. The different cDNA dones contain their respective cloned cDNA fragments which, on the whole, cover a region of at least the 333rd to 5177th nucleotides of the non-a, non-B hepatitis virus entire nucleotide sequence from the 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2(16) hereof. The cDNA fragments are taken out from the cDNA clones by cutting so as to respectively have predetermined nucleotide sequences such that when the predetermined nucleotide sequences are arranged in sequence, the resultant nucleotide sequence would have at least a region which coincides with the region of the 333rd to 5177th nucleotides.

The above-mentioned desired region of NANBV gene can be constructed by using, for example, BK 112, BK 146, BK 147, BK 157 and BK 166 selected from the NANBV cDNA clones disclosed in Fig. 1(1) and Fig. 1(2).

The taken-out cDNA fragments respectively having the above-mentioned predetermined nucleotide sequences are ligated in sequence to thereby construct a first deoxyribonucleic acid comprising a nucleotide sequence comprising at least the 333rd to 5177th nucleotides of the non-A, non-B hepatitis virus entire nucleotide sequence from 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2(16) hereof.

As the replicable expression vector which may be used in this step, any conventionally known or commercially available expression vector can be used. Examples of expression vectors include plasmid vector pSN508 for enterobacteria (U.S. Patent No. 4,703,005), plasmid vector pBH103 for yeast, and its series (Japanese Patent Application Laid-Open Specification No. 63-22098), plasmid vector pJM105 (Japanese Patent application Laid-Open Specification No. 62-286930), vaccinia virus WR strain (ATCC VR-119) and vaccinia virus LC16m8 strain (Japanese Patent Application Publication 55-23252), an attenuated varicella virus Oka strain (U.S. Patent No. 3,985,615), an attenuated Marek's disease virus [The Journal of Japanese Society of Veterinary, 27, 20-24 (1984), and Gan Monograph on Cancer Research, 10, 91-107 (1971)], plasmid vector pTTQ series (manufactured and sold by Amersham, England), plasmid vector pSLV series (manufactured and sold

by Pharmacia LKB, Sweden), and the like.

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The NANBV DNA-inserted expression plasmid vectors are individually introduced or transfected into host cells sensitive to the vector according to a conventional method, to obtain transformants which are capable of producing an NANBV particle. Then, the transformant(s) which has produced an NANBV particle is selected. The production of an NANBV particle may be detected by the immunoassay mentioned in Step (V). Further, the production of an NANBV particle may be confirmed or detected according to a conventional method, such as electron microscopy, immunoelectron microscopy, density-gradient centrifugation, light scattering photometry or the like. As mentioned above, when a plasmid is used as an expression vector, a transformant having a capability of producing NANBV particles may be obtained. On the other hand, when an animal virus gene is used as an expression vector, a recombinant virus which is capable of producing an NANBV particle is obtained.

By culturing the transformant or recombinant virus obtained above according to a customary method, an NANBV particle can be produced in the culture of the transformant or recombinant virus on a commercial scale. With respect to the details of the method in which an animal virus gene is used as an expression vector, reference may be made to European patent Application Publication No. 0 334 530 A1.

Accordingly, in still another aspect of the present invention, there is provided a method for producing an isolated non-A, non-B hepatitis virus particle, which comprises:

- (a) providing not more than ten different cDNA clones each comprising at least 1000 nucleotides and prepared from a non-A, non-B hepatitis virus genomic RNA fragment of at least 1000 nucleotides, said not more than ten different cDNA clones containing their respective cloned cDNA fragments which, on the whole, cover a region of at least the 333rd to 5177th nucleotides of the non-A, non-B hepatitis virus entire nucleotide sequence from the 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2(16) hereof;
- (b) taking out said cDNA fragments from said cDNA clones by cutting so as to respectively have predetermined nucleotide sequences such that when the predetermined nucleotide sequences are arranged in sequence, the resultant nucleotide sequence has at least a region which coincides with the region of the 333rd to 5177th nucleotides:
- (c) ligating said taken-out cDNA fragments respectively having said predetermined nucleotide sequences in sequence to thereby construct a first deoxyribonucleic acid comprising a nucleotide sequence comprising at least the 333rd to 5177th nucleotides of the non-A, non-B hepatitis virus entire nucleotide sequence from the 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2(16) hereof;
- (d) introducing at least one deoxyribonucleic acid selected from said first deoxyribonucleic acid and a second deoxyribonucleic acid obtained by substituting at least one nucleotide of the nucleotide sequence of said first deoxyribonucleic acid in accordance with the degeneracy of the genetic code into a replicable expression vector selected from a plasmid and an animal virus gene to obtain a replicable recombinant DNA comprising said plasmid and said at least one deoxyribonucleic acid introduced therein when said replicable expression vector is a plasmid or obtain a recombinant virus comprising said animal virus and said at least one deoxyribonucleic acid introduced therein when said replicable expression vector is an ani-
- (e) transfecting prokaryotic or eukaryotic cells with said recombinant DNA when said replicable expression vector used in step (d) is a plasmid, to thereby form a transformant, followed by selection of said transformant from parent cells of the prokaryotic or eukaryotic cell culture;
- (f) culturing said transformant obtained in step (e) in prokaryotic or eukaryotic cells to thereby produce a non-A, non-B hepatitis virus particle, or culturing said recombinant virus obtained in step (d) in eukaryotic cells to thereby produce a non-A, non-B hepatitis virus particle together with an animal virus; and
- (g) isolating said non-A, non-B hepatitis virus particle. In the above method, the deoxyribonucleic acid may preferably comprise a nucleotide sequence of the

333rd to 5918th nucleotides, a nucleotide sequence of the 333rd to 6371st nucleotides, a nucleotide sequence of the 333rd to 9362nd nucleotides, or a nucleotide sequence of the 1st to 9416th nucleotides.

It should be noted that in order to produce the NANBV particle of the present invention, the region of the NANBV cDNA to be expressed is required to contain all of the nucleotide sequences respectively coding for NS1 protein, NS2 protein and NS3 protein of NANBV in addition to all of the nucleotide sequences respectively coding for core antigen, matrix antigen and envelope antigen of NANBV.

Step (VII): Purification of an NANBV particle

The NANBV particle produced in the culture of the transformant or recombinant virus may be purified using an appropriate combination of customary techniques, for example, salting-out; adsorption and desorption using a silica gel, an activated carbon or the like; precipitation by an organic solvent; fractionation by ultracentrifugation; separation by ion exchange chromatography or affinity column chromatography; fractionation by highperformance liquid chromatography or electrophoresis, and the like.

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When the NANBV particle is purified from the culture of an E. coll transformant or a yeast transformant, from the viewpoint of effective removal of allergens derived from  $\underline{E}$ , coll and yeast which cause the quality of the produced NANBV particle to be markedly lowered, it is preferred that the purification be conducted by, for example, the steps of (1) adsorption and elution using a silica gel, removal of impurities by adsorption on an activated carbon and (2) fractionation by density gradient centrifugation in this order (Japanese Patent Application Laid-Open Specification No. 63-297). When the NANBV particle is purified from the culture of a recombinant virus, e.g., the culture of a recombinant virus-infected cells, a high purity NANBV particle can be obtained by subjecting a crude solution containing the particle to purification by ultracentrifugation and density gradient centrifugation repeatedly. Furthermore, for inactivating the NANBV particle in the culture to secure the safe handling of the particle and for fixing the particle to stabilize the immunogenicity and the antigenicity of the particle, it is preferred to add a conventional inactivating agent to the culture of the transformant or recombinant virus-infected cells, or to a culture liquid obtained by removing the transformant cells or the recombinant virusinfected cells. For example, an inactivating agent, such as formalin, may be added in an amount of from 0.0001 to 0.001 (v/v)% in the final concentration, followed by incubation at 4 to 37 °C for 5 to 90 days to thereby inactivate the NANBV particle. It should be noted that when an attenuated virus is used as an expression vector, an NANBV particle obtained from the recombinant virus can be used as an active ingredient for a live attenuated vaccine without the step of inactivation. The thus obtained NANBV particle suspension which is highly purified can be used for the preparation of a vaccine and a diagnostic reagent, as an original NANBV particle solution (an original NANBV vaccine solution).

In a further aspect of the present invention, there is provided a recombinant comprising a replicable expression vector selected from a plasmid and an animal virus gene and a deoxyribonucleic acid comprising at least one nucleotide sequence selected from the group consisting of a first nucleotide sequence of the 333rd to 5177th nucleotides of the non-A, non-B hepatitis virus entire nucleotide sequence from the 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2(16) hereof and a second nucleotide sequence obtained by substituting at least one nucleotide of said first nucleotide sequence in accordance with the degeneracy of the genetic code.

In the above-mentioned recombinant, the first nucleotide sequence may preferably comprise a nucleotide sequence of the 333rd to 5918th nucleotides, a nucleotide sequence of the 333rd to 6371st nucleotides, a nucleotide sequence of the 1st to 9416th nucleotides.

The replicable recombinant can be used not only for producing the NANBV particle of the present invention but also for amplifying the NANBV genomic cDNA to be used in the present invention by replication.

The purified NANBV particle of the present invention is useful as a diagnostic reagent for detecting NANB hepatitis.

The NANBV particle of the present invention can be formulated into a diagnostic reagent as follows. The purified NANBV particle solution obtained in Step (VII) mentioned above is dispensed in a vessel, such as a vial and an ampul, and sealed. The NANBV particle solution put in a vessel may be lyophilized before the sealing, in the same manner as mentioned above. The amount of the NANBV particle put in a vessel is generally about 1 µg to about 10 mg. Alternatively, the NANBV particle may also be adsorbed on the surface of a customarily employed support, such as a microplate, polystyrene beads, filter paper or a membrane.

The determination of the reactivity of the serum with the NANBV particle may be conducted in substantially the same manner as described in Step (V) mentioned above. That is, the determination of the reactivity may be conducted by a conventional immunoassay method, such as radioimmunoassay (RIA), enzyme-linked immunosorbent assay (ELISA), fluorescent antibody technique (FA), passive haemagglutination (PHA), reversed passive haemagglutination (rPHA) and the like. The amount of the NANBV particle to be used for the above immunoassay is generally from about 0.1 to about 100 mg/ml of serum. Particularly, the amounts of the NANBV particle to be used for RIA, ELISA, FA, PHA and rPHA are generally from 0.1 to 1 mg/ml, from 1 to 100 mg/ml, from 1 to 50 mg/ml and from 1 to 50 mg/ml, respectively.

The NANBV particle of the present invention may also be used for screening blood for transfusion. The screening method consists in:

- a) isolating serum from whole blood;
- b) contacting serum of an unknown blood with an isolated NANBV particle;
- c) determining whether the serum reacts with the NANBV particle;
- d) classifying the serum as positive or negative to non-A, non-B hepatitis based on the reactivity; and
- e) effecting separation of the blood in accordance with the identification.

The contact of serum of an unknown blood with the NANBV particle of the present invention, and the determination of the reactivity of the serum of the blood with the NANBV particle may be conducted in the same manner as mentioned above with respect to the method for diagnosing NANB hepatitis. By the above method,

a blood for transfusion free from the NANBV can be selected.

The polyclonal antibody and monoclonal antibody specific for the NANBV particle of the present invention may be used as an agent for removing NANBV from blood for transfusion. That is, NANBV present in blood can efficiently be removed by the polyclonal antibody or the monoclonal antibody by antigen-antibody reaction.

Further, the NANBV particle of the present invention may advantageously be used as an active ingredient of a vaccine for NANB hepatitis. The vaccine for NANB hepatitis may be prepared as follows. The culturing of a transformant containing a recombinant phage or plasmid carrying the cDNA coding for the NANBV particle, or a cell infected with the recombinant virus carrying the cDNA coding for the NANBV particle is conducted in the same manner as described above to thereby produce the NANBV particle in the culture. For inactivating the NANBV particle in the culture to secure the safety of the NANBV particle and for fixing the NANBV particle to stabilize the immunogenicity and the antigenicity of the particle, it is preferred to add a conventional inactivating agent to the culture of the transformant or recombinant virus-infected cell, or to a culture medium obtained by removing the transformant cells or the recombinant virus-infected cell. For example, an inactivating agent, such as formalln, may be added in an amount of from 0.0001 to 0.001 v/v%, followed by incubation at 4 to 37 °C for 5 to 90 days. Then, the resultant culture or culture medium is subjected to purification in the same manner as mentioned above. Thus, an original NANB hepatitis vaccine solution containing the purified NANBV particle is obtained.

The original NANBV hepatitis vaccine solution is filtered using a microfilter by a standard method to sterilize the solution. The filtrate is diluted with physiological saline so that the protein concentration is about 1 to about 500 µg/ml as measured by the Lowry method. Further, at least one stabilizing agent may be added. As the stabilizing agent, any commercially available stabilizing agent may be used. Examples of stabilizing agents include gelatin and hydrolysates thereof, human albumin, saccharides such as glucose, fructose, galactose, sucrose and lactose, and amino acids such as glycine, alanine, lysine, arginine and glutamine. Also, an adjuvant may be used to prepare an adsorbed vaccine. In this case, an adjuvant, such as, an aluminum hydroxide gel is added to the solution, before the addition of a stabilizing agent, so that the concentration of the added gel becomes about 0.1 to about 1.0 mg/ml, followed by mixing, thereby adsorbing the NANBV particle onto the adjuvant. As an adjuvant, there may also be employed precipitating depositary adjuvants such as calcium phosphate gel, aluminum phosphate gel, aluminum sulfate, alumina and bentonite, and adjuvants which are capable of inducing antibody production such as muramyl peptide derivatives, polynucleotides, Krestin® (manufactured and sold by Kureha Chemical Industry Co., Ltd., Japan) and picibanil (both of which are an antineoplastic agent).

Then, the thus obtained NANB hepatitis vaccine solution containing an (gel-adsorbed or non-adsorbed) NANBV particle is dispensed into a small vessel, such as an ampul and a vial, and sealed. Thus, there is obtained a purified (adsorbed or non-adsorbed) NANB hepatitis vaccine comprising an (adsorbed or non-adsorbed) NANBV particle.

The NANB hepatitis vaccine solution thus obtained may be lyophilized to obtain the NANB hepatitis vaccine in a dried form so that the product can be transported to and stored at a place of severe climate, for example, in an area in the tropics. The lyophilization may generally be conducted according to a standard method after the liquid (adsorbed or non adsorbed) NANB hepatitis vaccine is dispensed in a vessel such as a vial and an ampul. After lyophilization, a nitrogen gas is introduced in the vessel containing the dried vaccine, followed by sealing. Incidentally, the quality of the vaccine produced is examined in accordance with "Adsorbed Hepatitis B Vaccine", "Dried Japanese Encephalitis Vaccine", and "Adsorbed Pertussis Vaccine" provided for in Notification No. 159 of the Ministry of Health and Welfare, Japan, "Minimum Requirements for Biological Products".

The NANB hepatitis vaccine may be prepared in the form of a mixed vaccine which contains an adsorbed NANBV particle mentioned above and at least one antigen other than the present NANBV particle. As the antigen other than the present NANBV particle, there may be employed any antigens that are conventionally used as active ingredients of the corresponding vaccines insofar as the side effects and adverse reactions caused by such other antigens and the NANBV particle are not additively or synergistically increased by the use of the NANBV particle and such other antigens in combination and the antigenicities and immunogenicities of the NANBV particle and such other antigens are not reduced by the interference between the NANBV particle and other antigens. The number and the types of the antigens which may be mixed with the NANBV particle are not limited insofar as the side effects and adverse reactions are not increased additively or synergistically and the antigenicity and immunogenicity of each of the NANBV particle and such antigens are not reduced as mentioned above. Generally, two to six types of antigens may be mixed with the NANBV particle. Examples of antigens which may be mixed with the present NANBV particle, include detoxified antigens, inactivated antigens or toxoids which are derived from Japanese encephalitis virus, HFRS (hemorrhagic fever with renal syndrome) virus, influenza virus, parainfluenza virus, hepatitis B virus, dengue fever virus, AIDS virus, Bordetella pertussis, diphtheria bacillus, tetanus bacillus, meningococcus, pneumococcus and the like.

Generally, the vaccine comprising the NANBV particle of the present invention may be contained and sea-

led in a vial, an ampul or the like. The vaccine of the present invention may generally be administered in the form of a liquid or suspension. In the case where the vaccine is in a dried form, the vaccine is dissolved or suspended in sterilized distilled water before administration, the amount of the distilled water being such that the volume becomes the original volume before being subjected to lyophilization. Generally, the vaccine may be administered subcutane-ously. The dose of the vaccine may generally be about 0.5 ml. In general, the dose of the vaccine for a child may be half as much as that of the vaccine per adult. The vaccine may generally be administered twice at an interval of about one week to one month and then, about half a year later, administered once more.

Further, the NANBV particle may be used for preparing an antibody, such as a polyclonal antibody and a monoclonal antibody, specific for the NANBV particle. For example, a polyclonal antibody specific for the NANBV particle may be prepared by a conventional method as follows. The purified NANBV particle of the present invention is inoculated subcutaneously, intramuscularly, intraperitoneally or intravenously to an animal, such as mouse, guinea pig and rabbit. The inoculation of the NANBV particle is generally conducted several times at intervals of 1 to 4 weeks, to thereby completely immunize the animal. In order to enhance the immunizing effect, a conventional and commercially available adjuvant may be used. Then, blood serum is collected from the immunized animal and an anti-NANBV particle polyclonal antibody is isolated and purified from the blood serum according to a standard method.

On the other hand, a monoclonal antibody specific for the NANBV particle may be prepared by a conventional method as described, for example, in Cell Technology, 1, 23-29 (1982). For example, splenic cells obtained from a mouse immunized with the purified NANBV particle are fused with commercially available mouse myeloma cells by cell fusion technique, to obtain hybridomas. The hybridomas are screened to obtain a hybridoma capable of producing an antibody reactive with the NANBV particle. The obtained hybridoma is cultured in a standard method. From the supernatant of the culture, an anti-NANBV particle monoclonal antibody is isolated and purified by a standard method.

The above-mentioned polyclonal antibody and monoclonal antibody may also be used as a diagnostic reagent for diagnosing NANB hepatitis. The diagnosis of NANB hepatitis using the antibody may be conducted by immunoassay in substantially the same manner as mentioned above with respect to the diagnosis of NANB hepatitis using the NANBV particle. By the use of the polyclonal antibody or the monoclonal antibody, the identification and quantification of the NANBV particle present in a liver tissue and blood can be conducted.

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The NABV particle of the present invention has an extremely broad spectrum of antigenicity and specifically reacts with the serum not only of a chronic NANBV patient but also of an acute NANBV patient. Therefore, the NANBV particle is able to provide a diagnostic reagent of high reliability having not only a high detection ratio for an antibody but also a high precision in the detection. Further, when the NANBV particle of the present invention is used for screening blood for transfusion, blood which is infected by NANBV can be selected easily with high reliability and removed from blood not infected by NANBV. Therefore, the post-transfusion NANB hepatitis can be prevented.

Further, the NANBV particle of the present invention may advantageously be used as an active ingredient of a vaccine for preventing NANB hepatitis, which is extremely excellent in immunogenicity. In addition, a recombinant virus, e.g., recombinant vaccinia virus prepared by inserting the NANBV genomic cDNA into a vaccinia virus, is useful as an active ingredient of a vaccine.

Further, by the use of the NANBV particle of the present invention, an antibody, particularly monoclonal antibody, specific for NANBV can easily be prepared. The antibody specific for NANBV can advantageously be used as not only a diagnostic reagent for detecting NANB hepatitis, but also an agent for removing NANBV from blood for transfusion.

Furthermore, it should be noted that the NANBV particle of the present invention is not produced by the infection of an animal with a virus, but produced in an isolated form by gene expression of the DNA coding for the present NANBV particle in a host cell. Hence, the possibility of infection during the steps for production of the present NANBV particle is substantially eliminated. Also, the production cost can be decreased. Moreover, since all of the materials used in the production process, e.g., medium for the incubation system, are well-known in respect of the composition thereof, purification is facile and an NANBV particle product having high purity can be obtained.

By the present invention, it is possible to produce an isolated NANBV particle and its gene with high purity which cannot be found in nature. The produced NANBV particle and its gene can greatly contribute to researches on NANB hepatitis, hepatoma, liver cancer, etc.

The present invention will now be described in detail with reference to the following Examples and Reference Examples, which should not be construed to be limiting the scope of the present invention. Example 1 is divided into Part I and Part II, and Reference Examples 1-3 are inserted therebetween.

Example 1 (Part I)

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Step 1 (Preparation of a plasma-derived RNA for producing a cDNA, which is complementary to NANBV genome RNA)

In order to obtain NANBV from plasma, 4.8 liters of human plasma exhibiting a glutamic-pyruvic transaminase (GPT) activity of 35 IU/liter or more (as measured by the method of Wroblewski, F & J.S. LaDue: Serum glutamic-pyruvic transaminase in cardiac and hepatic disease. Proc. Soc. Exp. Biol. Med., 91:569, 1956) was applied on a 30 % (w/w) aqueous sucrose solution, and subjected to centrifugation at 48,000 x g at 4 °C and for 13 hours to obtain a precipitate. The precipitate was suspended in an aqueous solution containing 50 mM Tris-HCI (pH 8.0) and 1 mM EDTA, and once more subjected to centrifugation at 250,000 x g at 4 °C and for 3 hours to thereby obtain a precipitate. The obtained precipitate was dissolved in 75 ml of 5.5 M GTC solution containing 5.5 M quanidine thiocyanate, 20 Mm sodium citrate (pH 7.0), 0.05 % sarkosyl (sodium lauryl sarcosinate) and 0.1 M 2-mercaptoethanol. The resultant solution was applied on 16 ml of CsTFA-0.1 M EDTA solution (  $\rho$  = 1.51), and subjected to centrifugation at 140,000 x g at 15 °C and for 20 hours to thereby obtain a precipitate of RNA. The supernatant containing proteins and DNA was removed by suction, and the precipitate was dissolved in 200 μl of TE buffer solution containing 10 mM Tris-HCl (pH 8.0) and 1 mM EDTA. 20 μl of 3 M sodium chloride and ethanol were added to the solution, and allowed to stand still at -70 °C for 90 minutes. The mixture was centrifuged at 12,000 X g at 4 °C and for 30 minutes to obtain a precipitate. The precipitate was dissolved in TE, and sodium chloride and ethanol were added in the same manner as mentioned above. The mixture was allowed to stand still at -70 °C to obtain a precipitate. The precipitate was dissolved in 10 µl of TE to thereby obtain a purified RNA.

Step 2 (Preparation of a liver-derived RNA for producing a cDNA, which is complementary to NANBV genome RNA)

NANBV genome RNA was prepared from a liver tissue cut off from a NANBV hepatitis patient by the method of Okayama et al. (see H. Okayama, M. Kawaichi, M. Brown-stein, F. Lee, T. Yokota, and K. Arai: High-Efficiency Cloning of Full-Length cDNA; Construction and Screening of cDNA Expression Libraries for Mammalian Cells. Methods in Enzymology 154, 3-28, 1987).

Illustratively stated, 1 g of liver tissue was cut into small pieces. The small pieces were suspended in 100 ml of 5.5 M GTC solution as used in Step 1, and homogenized by means of a Teflon-glass homogenizer. Subsequently, the introduction of the homogenate into a syringe having #18 needle and the discharge of the homogenate from the syringe through the needle were repeated to thereby mechanically split DNA. The resultant homogenate was centrifuged at 1,500 x g (lower centrifugal force) at 4 °C and for 15 minutes to thereby obtain a supernatant. The supernatant was superposed on CsTFA solution and centrifuged in substantially the same manner as described in Step 1 to thereby obtain a precipitate as an RNA fraction. The thus obtained precipitate was suspended in 0.4 ml of 4 M GTC solution. 10 μl of 1 M acetic acid and 300 μl of ethanol were added to the suspension, and allowed to stand still at -20 °C for at least 3 hours to thereby obtain a precipitate of RNA. The precipitate was separated by centrifugation at 12,000 x g at 4 °C and for 10 minutes, and dissolved in 1 ml of TE solution. 100 μl of 2 M sodium chloride solution and 3 ml of ethanol were added to the solution, and the mixture was allowed to stand still at -20 °C for 3 hours. The resultant precipitate was collected by centrifugation and dissolved in 10 μl of TE to thereby obtain a purified, liver-derived RNA.

Step 3 (Preparation of a double-stranded cDNA using a cDNA synthesis kit)

A double-stranded cDNA was prepared using a commercially available cDNA synthesis kit (manufactured and sold by Amersham International, England).

Illustratively stated, 0.75  $\mu$ g of the purified RNA obtained in Step 1 and 2  $\mu$ l of random hexanucleotide primer and 2  $\mu$ l of reverse transcriptase taken from the reagents included in the kit were put in a reaction tube. Then, distilled water was added in an amount such that the total volume of the resultant mixture became 20  $\mu$ l. The mixture was incubated at 42 °C for 40 minutes, thereby preparing a first strand of cDNA. Subsequently, a second strand of cDNA was synthesized while cooling the reaction mixture in Ice water, as follows. To 20  $\mu$ l of the reaction mixture were added 37.5  $\mu$ l of buffer for second strand synthetic reaction, 1  $\mu$ l of E. C oil ribonuclease H and 6.8  $\mu$ l of DNA polymerase I, which were taken from the reagents included in the kit, followed by addition of 34.9  $\mu$ l of distilled water. The mixture was incubated at 12 °C for 60 minutes, 22 °C for 60 minutes and at 70 °C for 10 minutes. Then, the mixture was once more cooled with ice water. 1  $\mu$ l of T4 DNA polymerase was added, incubated at 37 °C for 10 minutes, and 4  $\mu$ l of 0.25 M EDTA (pH 8.0) was added to thereby terminate

the reaction. The reaction mixture was mixed well with a mixture of phenol and chloroform, and centrifuged at 12,000 x g for one minute to thereby separate an aqueous layer. The aqueous layer was again subjected to the same extraction as mentioned above, and an equal amount of chloroform was added. The mixture was agitated well and centrifuged to separate an aqueous layer. Subsequently, an equal amount of 4 M ammonium acetate and a two-fold amount of ethanol were added to the aqueous layer, and the mixture was cooled to -70 °C, thereby obtaining a precipitate of purified double-stranded cDNA. The precipitate was dissolved in 50 µl of 2 M ammonium acetate. To the mixture, 100 µl of ethanol was added, and the resultant mixture was cooled to -70 °C to thereby obtain a precipitate. The precipitate was collected by centrifugation at 12,000 x g for ten minutes. The collected precipitate was dried and then, dissolved in 20 µl of TE.

Step 4 (Preparation of a double-stranded cDNA by the Polymerase Chain Reaction (PCR) method)

The cDNAs which were prepared by means of a reverse transcriptase using as templates the RNAs prepared in Steps 1 and 2, were individually amplified by the PCR method (see Saiki, R. K., Gelfand, D. H., Stoffer, S., Scharf, S. J., Higuchi, R., Horn, G. T., Mullis, K. B., and Erlich, H. A., Primer-directed enzymatic amplification of DNA with a thermostable DNA Polymerase, Science 239:487-491, 1988). That is, 5 to 1,000 ng of the RNA was incubated in 20 μl of a reverse transcriptase solution containing 50 mM Tris-HCl (pH 8.3), 40 mM KCl, 6 mM MgCl<sub>2</sub>, 1 μM 3′-primer [synthesized oligonucleotide comprised of the 7949th to 7973rd 25 nucleotides in Fig. 2(14)], 10 mM dNTP, and 0.5 unit of reverse transcriptase (product of New England Bio Lab., U.S.A.) at 37 °C for 30 minutes. To the resultant mixture was added 80 μl of a PCR reaction solution containing 18 mM Tris-HCl (pH 8.3), 48 mM KCl, 1.5 mM MgCl<sub>2</sub>, 0.6 μM each of 5′-primer [synthesized oligonucleotide comprised of the 7612nd to 7636th 25 nucleotides in Fig. 2(13)] and the above-mentioned 3′-primer, 10 mM dNTP and 2.5 units of Taq DNA polymerase (manufactured and sold by Perkin Elmer Cetus Co., Ltd., U.S.A.). The mixture was subjected to incubation at 94 °C for one minute, at 50 °C for 2 minutes and at 72 °C for 3 minutes. This incubation was repeated 40 times. The resultant mixture was subjected to agarose gel electrophoresis, thereby obtaining amplified cDNA. The amplified cDNA was subjected tophenol treatment, ethanol precipitation and drying. The dried cDNA was dissolved in 10 μl of TE.

Step 5 (Preparation of a cDNA library using lambda gt11)

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Using a commercially available cDNA cloning kit (manufactured and sold by Amersham International, England), a cDNA library was prepared. That is, to 130 ng of cDNA prepared in step 3 were added 2  $\mu$ l of L/K buffer, 2  $\mu$ l of EcoRI adaptor and 2  $\mu$ l of T4 DNA ligase, which were taken from the reagents included in the cloning kit. Distilled water was added to the solution in an amount such that the total volume of the resultant mixture became 20  $\mu$ l. The mixture was incubated at 15 °C for 16 to 20 hours, and 2  $\mu$ l of 0.25 M EDTA was added thereto, to thereby terminate the reaction. Subsequently, the mixture was passed through a size fractionating column included in the kit, thereby removing EcoRI adaptors which were not ligated to the cDNA. To 700  $\mu$ l of the cDNA having EcoRI adaptor ligated thereto were added 83  $\mu$ l of L/K buffer and 8  $\mu$ l of T4 polynucleotidekinase. The mixture was incubated at 37 °C for 30 minutes. The resultant mixture was subjected to phenol extraction twice, concentration to 350 to 400  $\mu$ l by means of butanol and then ethanol precipitation, thereby obtaining a precipitate. The precipitate was dissolved in 5  $\mu$ l of TE.

Subsequently, in order to insert the cDNA having EcoRI adaptor ligated thereto to the EcoRI site of cloning vector lambda gt11, 1  $\mu$ I of L/K buffer, 2  $\mu$ I (1  $\mu$ g) of lambda gt11 arm DNA and 2  $\mu$ I of T4 DNA ligase were added to 1  $\mu$ I (10 ng) of the above-mentioned cDNA having EcoRI adaptor ligated thereto. Distilled water was added to the mixture in an amount such that the total volume of the mixture became 10  $\mu$ I. The mixture was incubated at 15 °C for 16 to 20 hours. Thus, a recombinant lambda gt11 DNA solution was prepared. Further, a recombinant lambda phage was obtained by in vitro packaging using a commercially available in vitro packaging kit (manufactured and sold by Stratagene Co., Ltd., U.S.A.) including Gigapack II Gold solutions A and B, SM buffer and chloroform. That is, 10  $\mu$ I of Gigapack II Gold solution A and 15  $\mu$ I of Gigapack II Gold solution B were added to 4  $\mu$ I of the above-mentioned recombinant lambda gt11 DNA solution. The mixture was incubated at 22 °C for 2 hours to obtain a recombinant phage. After the incubation, 470  $\mu$ I of SM buffer and 10  $\mu$ I of chloroform were added and the recombinant phage was stored at 4 °C.

Step 6 (Cloning of cDNA using E. coli Plasmid pUC19)

Using a commercially available DNA ligation kit (manufactured and sold by Takara Shuzo Co., Ltd., Japan) including solutions A and B, the cDNA was inserted in <u>E. coli</u> plasmid pUC19 (C. Yanishi-Perron, J. Vieira, J. Messing, Gene 33, 103, 1985), and cloned in <u>E. coli</u>. That is, 40 µl of solution A and 10 µl of solution B were

added to 5  $\mu$ l of the cDNA prepared by polymerase chain reaction (PCR) in Step 4 and 5  $\mu$ l (50 ng) of plasmid pUC19 DNA which had been digested with restriction enzyme Smal and dephosphorylated. The mixture was incubated at 15 °C for 16 hours. <u>E. coll</u> strain JM 109 (see Messing, J., Crea, R., and Seeburg, P.H., Nucleic Acids Res. 9, 309, 1981) was transformed with the above-obtained plasmid DNA according to the calcium chloride method (see Mandel, M. and A. Higa, J. Mol. Biol., 53, 154, 1970). Thus, a transformed <u>E. coll</u> containing the plasmid having the cDNA ligated thereto was obtained.

Step 7 (Screening of clone having NANBV gene from a cDNA library)

E. coli strain Y 1090 (see Richard A. Young and Ronald W. Davis, Science, 222, 778, 1983) was cultured in 50 ml of LBM medium containing 1 % tryptone, 0.5 % yeast extract, 1 % sodium chloride, 50 μg/ml ampicillin and 0.4 % maltose at 37 °C. The E. coli cells in a logarithmic growth phase were suspended in 15 ml of 10 mM magnesium sulfate cooled with ice. The bacteriophage solution obtained in Step 5 was diluted with SM buffer containing 0.1 M sodium chloride, 8 mM magnesium sulfate, 50 mM Tris-HCl (pH 7.5) and 0.01 % gelatin. 0.1 mi of the diluted phage solution was mixed with an equal volume of the above-mentioned E. coli cell suspension, and the mixture was incubated at 37 °C for 15 minutes. To the mixture was added 4 ml of soft agar medium containing 1 % tryptone, 0.5 % yeast extract, 0.5 % sodium chloride, 0.25 % magnesium sulfate and 0.7 % agar (pH 7.0) heated to 45 °C. The mixture was spread on L-agar plate containing 1 % tryptone, 0.5 % yeast extract, 1 % sodium chloride, 1.5 % agar and 100 μg/ml ampicillin (pH 7.0), and incubated at 42 °C for 3 hours. Subsequently, 10 mM IPTG (isopropyl β-D-thiogalactopyranoside) was infiltrated into a nitrocellulose filter and the nitrocellulose filter was dried and closely contacted with the plate. The plate in contact with the filter was incubated at 37 °C for 3 hours. The filter was separated, and washed with TBS buffer three times. The washed filter was immersed in 2 % bovine serum albumin solution, and incubated at room temperature for one hour. 1/20 volume of E. coli lysate solution included in a commercially available immunoscreening kit (manufactured and sold by Amersham International, England) was added to pooled serum from NANB hepatitis patients, and incubated at room temperature for 30 minutes. Thereafter, the serum was diluted to 50-fold with 0.2 % bovine serum albumin-added TBS buffer, and the filter was immersed in the diluted serum solution, and incubated at room temperature for one hour.

The resultant filter was washed four times with a TBS buffer containing 0.05 % Tween 20. The washed filter was immersed in an antibody solution which had been prepared by diluting a peroxidase-labeled anti-human IgG (manufactured and sold by Cappel Co., Ltd., Germany) 1,000-fold for one hour. The filter was washed with the above-mentioned Tween-TBS buffer, and immersed in a solution prepared by adding 0.4 ml of DAB (3,3'-diaminobenzidine tetrahydrochloride) and 15 µl of a 30 % aqueous hydrogen peroxide solution to 50 ml of a TBS buffer, followed by incubation at room temperature for 5 to 30 minutes to allow color development. The resultant filter was completely washed with distilled water to terminate the reaction.

By the above-mentioned procedure, the obtained plaques were purified. As a result, 9 positive clones were isolated, which were, respectively, designated as BK 102, BK 103, BK 105, BK 106, BK 108, BK 109, BK 110, BK 111 and BK 112. All of these clones did not react with serum from a healthy human, but reacted with serum from a patient suffering from NANB hepatitis. See Table 1.

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Table 1
Reactivity between the serum obtained from a patient suffering from NANB hepatitis and the recombinant lambda gtll phage clone

Clone	Serum from healthy person	Serum from NANB hepatitis patient
BK 102	0/10*	10/11
BK 103	0/10	9/11
BK 105	0/10	11/11
BK 106	0/10	11/11
BK 108	0/10	9/11
BK 109	0/10	9/11
BK 110	0/10	9/11
BK 111	0/10	9/11
BK 112	0/10	10/11

\* the number of positive samples/the number of specimens.

Step 8 (Determination of the nucleotide sequence of the obtained clones)

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Recombinant phage DNAs of clones BK 102 to BK 112 were purified, and the DNAs were digested with restriction enzyme EcoRI. Then, cDNA fragments of NANBV were isolated and the isolated cDNAs were individually inserted into plasmid pUC19 at EcoRI site. Using the plasmids, <u>E. Coli</u> strain JM 109 was transformed in substantially the same manner as in Step 6. Plasmid DNAs were obtained from the transformed <u>E. coli</u> and purified. The nucleotide sequence of each of the NANBV cDNAs was determined using 7-DEAZA sequencing kit (manufactured and sold by Takara Shuzo Co., Ltd., Japan; see Mizusawa, S., Nishimura, S. and Seela, F. Nucleic Acids Res., 14, 1319, 1986). The relationship between the nucleotide sequences of the obtained cDNA clones is shown in Fig. 1(1).

Step 9 (Cloning of NANBV cDNA clones from a cDNA library by Genomic Walking)

Probes were prepared by labeling with <sup>32</sup>P-dCTP the cDNA fragments of clone BK 102, clone BK 106 and clone BK 112 which were obtained in Step 8. Using the probes, phage clones containing NANBV cDNAs were obtained by hybridization from the cDNA library of cloning vector lambda gt11 obtained in Step 5. That is, plasmid DNAs were prepared from the transformed <u>E. coli</u> with clone BK 102, clone BK 106 and clone BK 112 obtained in Step 8 by the alkali method (see T. Maniatis, E.F. Fritsch, and J. Sambrook: Isolation of Bacteriophage λ and Plasmid DNA: "Molecular Cloning", Cold Spring Harbor Lab., pp 75-96.).

Plasmid DNA of clone BK 102 was digested with restriction enzymes Ncol and Hincil, and the resultant 0.7 kb fragments having been on the 5'-terminus side of the DNA were subjected to agarose gel electrophoresis, and collected. Plasmid DNAs of clone BK 106 and clone BK 112 were digested with restriction enzyme Ncol.

In the same manner as mentioned above, 1.1 kb DNA fragments were prepared from clone BK 106, and 0.7 kb fragments having been on the 3'-terminus side were prepared from clone BK 112. 25 ng to 1  $\mu$ g of DNA fragments were incubated with [ $\alpha$ -32P]dCTP (3000Cl/mmol; manufactured by Amersham Co., Ltd., England) at 37 °C for 3 to 5 hours, using commercially available DNA labeling kit (manufactured by Nippon Gene Co., Ltd.). Thus, probes for hybridization were prepared.

Subsequently, the cDNA library bacteriophage obtained in Step 5 was incubated at 42 °C in L-agar medium for 3 hours, as described in Step 7. Further, the bacteriophage was incubated at 37 °C for 3 hours, and was cooled. A nitrocellulose filter was contacted with phage plaques, and was allowed to stand still for 30 to 60 seconds. Thus, the phage plaques were adsorbed onto the filter.

The filter was subjected to alkali denaturation for 1 to 5 minutes using an aqueous solution containing 0.5 N sodium hydroxide and 1.5 M sodium chloride and to the neutralization with 0.5 M Tris-HCl (pH 8.0) containing 1.5 M sodium chloride for 1 to 5 minutes. The filter was washed with 2 x SSC solution containing 0.3 M sodium chloride and 0.03 M sodium citrate, dried, and baked at 80 °C for 2 hours.

The filter was incubated at 42 °C for 6 hours in a solution for hybridization containing 50 % formamide, 5 x SSC, 5 x Denhart solution, 50 mM phosphoric acidetric acid buffer (pH 6.5), 100  $\mu$ g/ml salmon sperm DNA and 0.1 % SDS. Then, the filter was Immersed in 300 ml of the hybridization solution having 1 ml of the above-mentioned probe of about 4 x 108 cpm/ml added thereto, and incubated at 42 °C for 16 to 20 hours. The filter was washed with a 2 x SSC solution containing 0.1 % (w/w) SDS four times and with a 0.1 x SSC solution containing 0.1 % (w/w) SDS twice. After the washing, the filter was dried, and was subjected to autoradiography. Thus, hybridization positive clones were isolated. As a result, 27 clones being reactive with the probe derived from clone BK 102, 14 clones being reactive with the probe derived from clone BK 106 and 13 clones being reactive with the probe derived from clone BK 106 and 13 clones being reactive with the probe derived from clone BK 108 and 13 clones being reactive with the probe derived from clone BK 108 and 13 clones being reactive with the probe derived from clone BK 108 and 13 clones BK 114 to BK 169.

The nucleotide sequence of each of clones BK 114 to BK 169 was determined according to the method described in Step 8, followed by mapping for each of the clones. As a result, a map of nucleotide sequence having a length of about 9.5 kb considered to be the approximately total length of the NANBV genome was obtained [see Fig. 1(2)].

Clone BK 157 located on the 5' terminus side was digested with restriction enzyme KpnI to thereby isolate a 0.55 kb fragment having been on the 5'-terminus side. Also, clone BK 116 located on the extreme 3'-terminus side was digested with restriction enzymes HpaI and EcoRI to thereby isolate a 0.55 kb fragment having been on the 3'-terminus side. A probe labeled with <sup>32</sup>P was prepared in the same manner as described above, and the cDNA library bacterio phageobtained in Step 5 was subjected to plaque hybridization. As a result, three new additional clones were separated by the probe derived from the clone BK 157. These new clones were, respectively, designated as clones BK 170, BK 171 and BK 172.

Step 10 (Analysis of the nucleotide sequence of cDNA)

The entire nucleotide sequence of NANBV gene was determined from the nucleotide sequences of the clones obtained in Steps 8 and 9, and shown in Figs. 2(1) to 2(16). From the Figures, it was assumed that the cloned genomic cDNAs of NANBV were composed of 9416 nucleotides, wherein there was an open reading frame composed of 9030 nucleotides coding for a protein composed of 3010 amino acid residues. The hydrophilicity/hydrophobicity pattern of this protein was similar to that of flavivirus as already reported (see H. Sumiyoshi, C. Mori, I. Fuke et al., Complete Nucleotide Sequence of the Japanese Encephalitis Virus Genome RNA. Virology, 161, 497-510, 1987). Clone BK 157 covers nucleotide numbers 1 to 1962 of Figs. 2(1) to 2(16), clone BK 172 covers nucleotide numbers 5 to 366, clone BK 153 covers nucleotide numbers 338 to 1802, clone BK 138 covers nucleotide numbers 1755 to 5124, clone BK 129 covers nucleotide numbers 4104 to 6973, clone BK 108 covers nucleotide numbers 6886 to 8344 and clone BK 166 covers nucleotide numbers 8082 to 9416. They are preserved as Escherichia coll BK 108 (deposited at Fermentation Research Institute, Japan under accession number FERM BP-2971), BK 129 (deposited at Fermentation Research Institute, Japan under accession number FERM BP-2972), BK 138 (deposited at Fermentation Research Institute, Japan under accession number FERM BP-2973), BK 153 (deposited at Fermentation Research Institute, Japan under accession number FERM BP-2974), BK 157 (deposited at Fermentation Research Institute, Japan under accession number FERM BP-3243), BK 166 (deposited at Fermentation Research Institute, Japan under accession number FERM BP-2975), and BK 172 (deposited at Fermentation Research Institute, Japan under accession number FERM BP-2976), respectively.

#### Reference Example 1

(Production of NANBV-related antigens in  $\underline{E}$ ,  $\underline{coll}$ , which antigens are related with the antibody-response accompanying NANBV infection)

Respective cDNAs of clone BK 106, clone BK 111 and clone BK 112 each obtained in Step 8 of Example 1 (Part I) and cDNA of clone BK 147 obtained in Step 9 of Example 1 (Part I) were individually inserted into plasmids, and the thus obtained plasmid DNAs were collected by the conventional alkali method. Subsequently, the collected DNA of clone BK 106 was digested with restriction enzymes EcoRI and ClaI to thereby obtain 0.5 μg of a DNA fragment of 0.34 kb in length. The thus obtained DNA fragment was incubated at 37 °C for 60 minutes in a T4 DNA polymerase solution containing 67 mM Tris-HCl (pH 8.8), 6.7 mM magnesium chloride, 16.6 mM ammonium sulfate, 10 mM 2-mercaptoethanol, 6.7 μM EDTA, 0.02 % bovine serum albumin, 0.3 mM dNTP and 2-5 units of T4 DNA polymerase, thereby rendering both terminals blunt. The DNA of clone BK 102 was digested with restriction enzyme BamHI to thereby obtain 0.5 μg of a DNA fragment of 0.7 kb in length, and the terminals of the DNA fragment were rendered blunt using T4 DNA polymerase in substantially the same manner as mentioned above. The DNA of clone BK 147 was digested with restriction enzyme Sau3Al to thereby obtain 0.5 µg of a DNA fragment of 1 kb in length and the terminals of the DNA fragment were rendered blunt in the same manner as mentioned above. Also, the DNA of clone BK 111 was digested with restriction enzyme EcoRI to thereby obtain 0.5 μg of a DNA fragment of 1 kb in length, and the terminals of the DNA fragment were rendered blunt in substantially the same manner as mentioned above. Subsequently, the DNA of expression vector pKK 233-2 (Amann, E. and J. Brosius. ATG vector for regulated high-level expression of cloned genes in Escherichia coli. Gene, Vol. 40, 183, 1985) was digested with restriction enzyme Hindlil. 2 µg of the resultant DNA was incubated at 37 °C for 20 minutes in a S1 nuclease solution containing 0.3 M sodium chloride, 50 mM sodium acetate (pH 4.5), 1 mM zinc sulfate and 100-200 units of S1 nuclease, and the reaction was terminated by adding 1/10 volume of each of 0.12 M EDTA and 1 M Tris-HCl solution (pH 9.0). Then, phenol extraction was performed, and the vector DNA having blunt terminals was precipitated by ethanol and collected. On the other hand, the DNA of vector pKK 233-2 was digested with restriction enzyme Pstl, and the digested DNA was purified by extraction with phenol and precipitation from ethanol. The terminals of 2 µg of the purified vector DNA which had been cleaved by restriction enzyme Psti were rendered blunt by the above-mentioned T4 DNA polymerase reaction. The thus obtained DNA fragments derived from clone BK 106 and clone BK 111 were each cleaved with restriction enzyme Hindill. 0.5 µg of each of the cleaved DNA fragments was mixed with 0.5 μg of a vector DNA having blunt terminals. The DNA fragments derived from clone BK 102 and clone BK 147 were each cleaved with restriction enzyme Pstl. 0.5 μg of each of the cleaved DNA fragments was mixed with 0.5 µg of a vector DNA having terminals thereof rendered blunt. The volume of each of the mixtures was adjusted to 20 μl by adding 2 μl of 10 x ligation solution containing 500 mM Tris-HCl (pH 7.5), 100 mM magnesium chloride, 100 mM DTT and 10 mM ATP, 300-400 units of T4 DNA ligase and distilled water. The mixtures were incubated at 14 °C for 12-18 hours, thereby obtaining plasmids, which were respectively designated as pCE-06, pE-11, pB-02 and pS-09. Using each of these plasmid DNAs, E. coli strain JM 109 was transformed in substantially the same manner as described in Step 6 of Example 1 (Part I), thereby obtaining transformed E. coli. The transformed E. coli was cultured at 37 °C in LB medium (pH 7.5) containing 1 (w/v)% trypton, 0.5 (w/v)% yeast extract and 1 (w/v)% sodium chloride, and when it was in logarithmic growth phase, 1 mM IPTG (Isopropyl-β-D-thiogalactopyranoside) was added to the medium. The culturing was further continued for 3 hours. Then, E. coli cells were collected by centrifugation (10,000 x g for 15 minutes), and the collected cells were lysed in 50 mM Tris-HCl (pH 8.0). The mixture was subjected to ultrasonic treatment (20 KHz, 600 W, 5 minutes), and centrifuged at 10,000 x g for 15 minutes to thereby obtain a supernatant fraction and a precipitate fraction. Each of the fractions was dissolved in a sample buffer containing of 20 (v/v)% glycerol, 0.1 M Tris-HCl (pH 6.8), 2 (w/v)% SDS, 2 (v/v)% 2-mercaptoethanol and 0.02 % BPB, heated at 100 °C for 3 minutes, and subjected to electrophoresis using 0.1 % SDS-7.5 % polyacrylamide gel to separate protein. After the electrophoresis, the protein was transferred to a nitrocellulose filter by trans blot cell (manufactured and sold by BIO-RAD Co., Ltd., U.S.A.). The filter was immersed in 3 % gelatin solution, and allowed to stand still for 60 minutes. The filter was incubated together with serum from a patient suffering from NANB hepatitls, which had been diluted 100-fold, for 2 to 3 hours at room temperature. The filter was washed with distilled water and then with TTBS solution containing 0.02 M Tris-HCl (pH 7.5), 0.5 M sodium chloride and 0.05 (v/v)% Tween 20. Subsequently, the washed filter was immersed in a 2,000 fold-diluted solution of peroxidase-labeled antihuman IgG antibody, and incubated at room temperature for 90 minutes. The filter was washed with distilled water and then with TTBS solution. The washed filter was immersed in a buffer having, added thereto, coloring agent DAB and 30 %, based on substrate, hydrogen peroxide as described in Step 7 of Example 1 (Part I) for 5 to 30 minutes, following by washing with water, to terminate the reaction.

As a result, as shown in Table 2, all of the antigens produced by the plasmids specifically react with serum

from a patient suffering from NANB hepatitis, thereby demonstrating that the proteins produced by the cDNAs inserted in the plasmids are clinically important.

Table 2
Reactivity evaluated by the Western blot method
between proteins produced by various plasmids and
sera from a patient suffering from NANB hepatitis

Plasmid	origin of	Extract	Serum from NANB hepati- titis patient	Serum from healthy human
pCE-066	BK 106	s	±	-
-		P	+	-
pE-11-89	BK 111	s	±	•
-		P	+	-
pB-02-10	BK 102	S	+	· <b></b>
•	•	P	••	
ps-09-07	BK 109	s	±	-
_		<b>P</b> .	+	-
pKK233-3	-	s	-	_
_		P	<b></b>	_

- S: Supernatant by centrifugation
- P: Precipitate by centrifugation
- +: positive
- t: slightly positive
- -: negative

#### Reference Example 2

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(Purification of NANBV-related antigens produced by E. coli and reactivity thereof with serum from a patient suffering from hepatitis)

The usefulness of the protein produced by the cDNA which was inserted into an expression vector was demonstrated by purifying the protein and using the purified protein as an antigen for ELISA or radioimmunoassay. That is, the lysate of the transformed E. coli which was obtained in Reference Example 1 was subjected to centrifugation at 10,000 x g for 15 minutes, thereby obtaining a supernatant and a precipitate. For example, the precipitate obtained from transformant JM 109/pCE 066 was suspended in a solution of 100 mM Tris-HCl (pH 8.0) and 0.1 % Triton X-100, and the resultant suspension was subjected to ultrasonic treatment at a frequency of 20 KHz (600 W) for one minute, followed by centrifugation at 21,000 x g for 15 minutes, thereby obtaining a precipitate. The precipitate was re-suspended in a solution of 100 mM Tris-HCl (pH 8.0) and 6 M urea, and then subjected to ultrasonic treatment followed by centrifugation.

The resultant supernatant was dialyzed against a solution of 10 mM phosphate buffer (pH 7.5) and 6 M urea to thereby obtain an antigen solution. 20 ml of the antigen solution was passed through a column (21.5 x 250 mm) packed with hydroxyapatite, which had been equilibrated with the above-mentioned buffer, to cause

the antigen to be adsorbed onto the packing material. The column was subjected to high speed liquid chromatography (HPLC) wherein elution was performed with the above-mentioned buffer having, added thereto, sodium chloride, the concentration of which was varied from 0 to 2 M with a linear concentration gradient, thereby obtaining a fraction containing an antigen. The obtained fraction was dialyzed against 50 mM carbonate buffer (pH 9.6) containing 0.05 % sodium dodecyl sulfate (SDS).

Further, the supernatant obtained by centrifugation (at 10,000 g for 15 minutes) of the lysate of transformant JM 109/pB-02-10 was treated with 35 % saturated ammonium sulfate, and the obtained precipitate was dissolved in 50 mM Tris-HCl (pH 8.5) buffer containing 100 mM 2-mercaptoethanol. The resultant solution was dialyzed against the above-mentioned buffer. Subsequently, 100 ml of the dialysed solution was passed through a column (22.0 x 200 mm) packed with DEAE cellulose, which had been equilibrated with the above-mentioned buffer, to cause the antigen to be adsorbed onto the packing material. The column was subjected to high performance liquid chromatography wherein elution was performed with 50 mM Tris-HCl (pH 8.5) buffer containing 100 mM 2-mercaptoethanol having, added thereto, sodium chloride, the concentration of which was varied from 0 to 2 M with a linear concentration gradient, thereby pooling a fraction containing the antigen.

The fraction was dialyzed against a solution of 10 mM phosphate buffer (pH 6.8) and 100 mM 2-mercaptoethanol. The dialyzed solution was passed through the column of hydroxyapatite for high performance liquid chromatography, which had been equilibrated by the above-mentioned buffer, to cause the antigen to be adsorbed onto the packing material. The column was subjected to high performance liquid chromatography wherein elution was performed with phosphoric acid, the concentration of which was varied with a linear concentration gradient from 10 to 400 mM, thereby pooling a fraction containing the antigen. The resultant fraction was dialyzed against 50 mM carbonate buffer (pH 9.6) containing 0.05 % SDS.

The precipitate obtained by centrifugation of the lysate of transformant JM 109/pE-11-89 was suspended in 10 mM phosphate buffer (pH 5.5). The suspension was subjected to the above-mentioned ultrasonic treatment for one minute, and then subjected to centrifugation at 21,000 x g for 15 minutes. The resultant precipitate was suspended in 100 mM carbonate buffer (pH 10.5) containing 500 mM sodium chloride and 10 mM EDTA. The resultant suspension was again subjected to the ultrasonic treatment for one minute, followed by centrifugation. The resultant supernatant was dialyzed against 30 mM phosphate buffer containing 6 M urea. Subsequently, 20 ml of the dialyzed solution was passed through a CM cellulose column (22 x 200 mm) for high performance liquid chromatography (HPLC), which had been equilibrated with the same buffer as used for the above-mentioned dialysis, to thereby cause the antigen to be adsorbed onto the packing material. The column was subjected to high performance liquid chromatography wherein elution was performed with the above-mentioned buffer having, added thereto, sodium chloride, the concentration of which was varied from 0 to 1.5 M with a linear concentration gradient, obtaining a fraction containing the antigen. The fraction was dialyzed against 50 mM carbonate buffer (pH 9.6) containing 0.05 % SDS, thereby obtaining a solution containing the antigen.

The antigens prepared above were used as an antigen for ELISA for the clinical diagnosis of infection with non-A, non-B hepatitis virus. That is, the protein concentration of each of the above-mentioned purified antigens was adjusted to 1 µg/ml, and put in each well of Microplate Immulone 600 (manufactured and sold by Greiner. Co., Ltd., Germany) in an amount of 100 µl for use in ELISA, which well was allowed to stand still at 4 °C overnight. The contents of the individual wells were washed well three times with PBS-T buffer containing 10 mM phosphate buffer (pH 7.2), 0.8 % sodium chloride and 0.05 % Tween 20, and sample serum diluted with the PBS-T buffer was added in an amount of 100 µl/well, followed by reaction at 37 °C for one hour. The contents of the individual wells were washed three times with the PBS-T buffer, and a peroxidase-labeled anti-human lgG antibody (manufactured and sold by Cappel Co., Ltd., Germany) which had been diluted 8000-fold with PBS-T buffer containing 10 % fetal calf serum was added in an amount of 100 µl/well. The individual well contents were reacted at 37 °C for one hour, and washed with the PBS-T buffer four times. A substrate coloring agent solution composed of 9 ml of 0.05 M citric acid-phosphate buffer and, contained therein, 0.5 µg of o-phenylenediamine and 20 µl of aqueous hydrogen peroxide, was added in an amount of 100 µl/well. The plate was light shielded, and allowed to stand still at room temperature for 60 minutes. 75 µl of 4 N sulfuric acid was added to each of the wells, and the absorbance at 490 nm was determined. The results are shown in Table 3. As apparent from the table, all of the antigens derived from the transformants specifically react with the serum from NANB hepatitis patient, thereby attesting to the usefulness in clinical diagnosis of the antigens produced by the transofrmants.

#### Table 3

Reactivity in ELISA between the purified antigens from various transformed <u>Escherichia coli</u> and the serum from NANB hepatitis patient

Serum from blood transfused patient of hepatitis

origin of antigen (transformed healthy hepato-Escherichia human coli) acute chronic cirrhosis hepatoma serum JM109/pCE-066 2/3\* 7/8 3/3 3/4 0/10 JM109/pB-02-10 2/3 8/8 4/4 3/3 0/10 JM109/pE-11-B9 2/3 8/8 2/4 3/3 0/10

\*: the number of positive samples/the number of samples examined

The same results as shown in Table 3 were also obtained by radioimmunoassay using the above-mentioned antigens. That is, a polystyrene bead of 1/4 inch in diameter (manufactured and sold by Pesel Co., Ltd., Germany) was put in 0.2 ml of each of the above-mentioned purified antigen solutions of 1 µg/ml in concentration, and allowed to stand still at 4 °C overnight. Then, the polystyrene bead was washed five times with the same PBS-T buffer as used in the above-mentioned ELISA, and a sample serum diluted 20 to 2500-fold with the PBS-T buffer was added in an amount of 200 µl/bead. Reaction was performed at 37 °C for 60 min. The polystyrene bead was washed five times with the PBS-T buffer, and 1251-labeled anti-human IgG antibody was added in an amount of 200 µl/bead. Reaction was performed at 37 °C for one hour and the bead was washed five times with the PBS-T buffer. The cpm of 1251 bound to the polystyrene bead was measured, thereby obtaining the same results as shown in Table 3. Thus, the usefulness of the purified antigens obtained above in the clinical diagnosis of infection with NANB hepatitis virus, was demonstrated.

#### Reference Example 3

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[Detection of NANBV nucleic acid according to PCR (Polymerase Chain Reaction) method]

For preventing NANB hepatitis caused by blood transfusion, it is important to determine whether or not any NANBV infection exists in the blood supplied for transfusion. Further, for diagnosing hepatitis, it is extremely clinically important to study whether or not any NANBV infection exists in liver tissue. The NANBV cDNA obtained according to the present invention can be advantageously used for producing a primer for polymerase chain reaction (PCR) useful for detecting NANB hepatitis. That is, as described in Step 1 of Example 1 (Part I), the purification of RNA was performed from 1 ml of each of sera derived from a patient and a healthy human. Likewise, RNA was prepared from liver cells as described in Step 2 of Example 1 (Part I). Subsequently, as described in Step 4 of Example 1 (Part I), PCR and electrophoresis were conducted to thereby prepare cDNAs. According to the customary procedure, whether or not the amplified cDNA was derived from NANBV, was investigated by Southern hybridization using 32P-labeled probe prepared from the cDNA derived from NANBV cDNA clone BK 108.

The results are shown in Table 4. From the table, it is apparent that the NANBV nucleic acid in serum can be detected and the serum infection with NANBV can be diagnosed by the use of the primer prepared from the nucleotide sequence of the NANBV cDNA obtained according to the present invention and the fragment of

cloned NANBV cDNA as a probe.

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Table 4

Detection of NANBV nucleic acid by PCR

sample		antibody against NANBV	PCR
serum from chroni	c hepatitis pat	tient	
NANB	1	+	+
	2	+	+
HBV carrier	1	-	
	2	-	-
healthy human	1 2	-	_
_	2	· <b>-</b>	-
excised liver		+	
from NANB hepatom			
cancerous si	te		+
non-cancerou	s site		+
excised liver		+	
from NANB hepatom			
cancerous si	te		+
non-cancerou	s site		+

Example 1 (Part II)

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step 1 (Construction of the plasmids for the expression of the entire coding region of the NANBV genomic cDNA in E. coli)

cDNA was isolated from each of clones BK112, BK146, BK147, BK157 and BK166 shown in Fig. 1 (1) and Fig. 1 (2), and plasmids for the expression of the entire coding region of the NANBV gene in <u>E. coli</u> were prepared as follows.

The plasmid DNA of clone BK157 was digested with restriction enzyme BamHI and subjected to agarose gel electrophoresis to thereby obtain a DNA fragment of 1.3 kb in length. The DNA fragment was inserted in plasmid pUC19 (manufactured and sold by Takara Shuzo Co., Ltd., Japan) at its BamHI site to thereby obtain plasmid pBam157. The plasmid pBam157 was digested with restriction enzymes Xbal and Ncol to thereby obtain a DNA fragment of about 3.9 kb in length. Separately, an oligonucleotide of 93bp (having 4 nucleotides deleted from the 3'-terminus side) was synthesized by ligating the sequence of the promoter region of 20bp (TAATAC-GACTCACTATAGGG) of bacteriophage T7 RNA polymerase having, attached thereto, Xbal linker sequence to the sequence of nucleotide numbers 1 to 73 shown in Fig. 2(1) having, attached thereto, Ncol linker sequence. The thus obtained oligonucleotide was ligated to the above-mentioned DNA fragment of 3.9 kb, thereby obtaining plasmid pDM-16. Then, pDM-16 was digested with restriction enzymes Clal and EcoRI to obtain a DNA fragment of about 3.5 kb. Separately, the DNA of clone BK146 was digested with restriction enzymes Cial and EcoRI to obtain a DNA fragment of 4.1 kb in length. The above-mentioned DNA fragment of about 3.5 kb was ligated to the thus obtained DNA fragment of 4.1 kb, to thereby obtain plasmid pDM-9. Then, plasmid pDM-9 was digested with SacII, thereby obtaining a DNA fragment of 2.7 kb and a DNA fragment of 4.9 kb. The DNA fragment of 4.9 kb was ligated at its SacII site with T4 DNA ligase and then digested with BamHI and EcoRI, thereby obtaining a DNA fragment of about 7.5 kb. Separately, the DNA of clone BK147 was digested with BamHI and EcoRI, thereby obtaining a DNA fragment of about 2 kb. The thus obtained DNA fragment was ligated to the above-mentioned DNA fragment of 7.5 kb to thereby obtain plasmid pBE147. Plas-

mid pBE147 was digested with Sacil. The above-mentioned DNA fragment of 2.7 kb derived from pDM-9 was inserted into the Sacil-digested pBE147, thereby obtaining plasmid pDM-B3. Plasmid pDM-B3 was digested with Xbal and EcoRI to thereby obtain a DNA fragment of 6.7 kb.

The DNA of clone BK166 was digested with BamHI to obtain a DNA fragment of 1.3 kb. This fragment was inserted in pUC19 at its BamHI site to obtain pBam166. pBam166 was digested with Ndel and HindIII to thereby obtain a DNA fragment of 2.8 kb. The DNA of clone BK112 was digested with EcoRI and Ndel to obtain a DNA fragment of about 1.6 kb. pUC19 was digested with EcoRI and HindIII to obtain a DNA fragment of about 2.6 kb. The above-obtained three types of DNA fragments were mixed and reacted with T4 DNA ligase to thereby obtain plasmid pEN112 in which these fragments were ligated together at their EcoRI site, Ndel site and HindIII site. Plasmid pEN112 was digested with EcoRI and Xbal to obtain a DNA fragment of 2.7 kb. pDM-B3 was digested with EcoRI and Xbal to obtain a fragment of 6.7 kb. The above-mentioned fragment of 2.7 kb was ligated to the fragment of 6.7 kb, and the resultant ligated DNA fragment was inserted in pUC19 at its Xbal site, thereby obtaining plasmids pDM-22 and pDM-18. Plasmid pDM-18 can be used for the transformation of an animal cell or the like so that the cell can produce an NANBV particle. The transformation can also be performed using an RNA prepared by transcripting pDM-18 by means of in vitro Transcription Kit (manufactured and sold by Boehringer Mannheim Yamanouchi, Japan). Plasmid pDM-22, in which the cDNA was Inserted in an orientation opposite to that of pDM-18, was digested with HindIII and Clal to obtain a DNA fragment of about 9 kb.

The DNA of clone BK106 was digested with BamHI to obtain a DNA fragment of about 1.0 kb. This fragment was inserted in plasmid pUC19 at its BamHI site to obtain plasmid pBam108. The thus obtained plasmid DNA was digested with Ncol, and the sticky terminus was rendered blunt with Mang Bean nuclease (manufactured and sold by Takara Shuzo Co., Ltd., Japan). The resultant plasmid was further digested with Xbal, thereby obtaining a fragment of about 3.6 kb. A synthetic oligonucleotide prepared by ligating the sequence of nucleotide numbers 333 to 372 shown in Fig. 2(1) to the downstream of Xbal linker, was ligated to this fragment to thereby obtain plasmid pXb106. Plasmid pXb106 was digested with HindliI and Clai to obtain a DNA fragment of 0.4 kb. This fragment was ligated to the above-mentioned fragment of about 9 kb derived from plasmid pDM-22 to thereby obtain plasmid pORF-24. Plasmid pORF-24 was digested with Xbal to obtain a DNA fragment of about 9.0 kb. This fragment was ligated to an expression vector (see F. William Studier and B.A.Moffatt, J. Mol. Blol., 189, 113, 1986) having, ligated thereto, T7 RNA polymerase gene promotor, thereby obtaining expression plasmid pJF-22.

Step 2 (Preparation of transformant E. coli and culturing thereof)

Using expression plasmid pJF-22 constructed in Step 1, <u>Escherichia coli</u> strain JM109 (DE3) (manufactured and sold by Promega Co., U.S.A.) was transformed by the calcium chloride method ( Journal of Molecular Biology, 53, 154, 1970), thereby obtaining trans formant JM109 (DE3) /pJF-22.

Transformant E. coli JM109 (DE3)/pJF-22 was subjected to the subsequent procedure as described in Reference Example 1. That is, the E. coli was cultured on LB culture medium, and then 0.5 mM IPTG was added thereto, followed by further culturing for 3 hours. After that period, the cultured cells were collected and heated in a buffer containing 2 % SDS and 2 % 2-mercaptoethanol at 100 °C for 3 minutes. The resultant cells were subjected to electrophoresis using a gel containing 0.1 (w/v)% SDS and 12.5 (w/v)% acrylamide. The resultant protein isolated on the gel was blotted onto a nitrocellulose membrane by means of a trans blot apparatus (manufactured and sold by Nippon Eido Co., Ltd., Japan) and subjected to Western blotting analysis to identify the obtained protein. In the Western blotting analysis, the specific antisera was used which was obtained by purifying the NANBV-related antigen prepared from the transformant obtained in Reference Example 1 and immunizing guinea pigs therewith. As a result of the Western blotting analysis, it was found that the protein produced by transformant JM109 (DE3)/pJF-22 reacted with all of the antisera (see Table 5).

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#### Table 5

Reactivity of protein produced in transformant

E. coli JM109 (DE3)/pJF-22 with NANBV-related
antibody

10			from litis pa		guinea	pig ant	ig antiserum	
	cell extract	pooled serum	acute	chronic	anti- çore	anti- NS3	anti- NS5	
15	JM109(DE3)/ PJF22	+*	+	+	+	+	+	
20	JM109(DE3)	-	-	-	-	-	-	

 Reactivity as measured by Western blotting analysis.

Thus, it was demonstrated that in this transformant, the expression was attained of the entire coding region of the NANBV gene from the 5'-end of the genome coding for the core antigen through the 3'-end of the genome coding for the non-structural protein NS5. From the results, it is apparent that this transformant can provide an antigen which is extremely useful for producing not only a diagnostic reagent for NANBV infection but also a vaccine for NANBV.

Step 3 (Production of NANBV particles by expression of NANBV genomic cDNA in animal cells)

Plasmid pORF-24 obtained in Step 1 was partially digested with Xbal and subjected to low melting point agarose gel electrophoresis to obtain a DNA fragment of about 9 kb in length. The thus obtained DNA fragment was inserted into plasmid pMAM-neo (available from Clontech, U.S.A.) which had been cleaved with Nhel, to thereby construct expression plasmid pMAM-neo10. The expression plasmid was transfected into cells of human hepatocyte Chang Liver (ATCC CCL 13) and Chimpanzee hepatocyte (purchased from Dalnippon Pharmaceutical Company, Ltd., Japan) by a calcium phosphate method ("Molecular Cloning", 16.33-16.39, Cold Spring Harbor Laboratory, 1989). The hepatocyte cells having plasmid pMAM-neo10 introduced thereto were rendered resistant to aminoglycoside antibiotic G418 so that the cells were able to form colonies in the presence of G418 in an amount of 600 µg/ml. Utilizing this resistance as a criterion, transformants were selected, followed by cloning. Transformant clones HL-A1 and HL-A2 produced from human hepatocyte and transformant clones CL-B11 and CL-B14 produced from chimpanzee hepatocyte were individually cultured in Eagle's MEM medium having, incorporated therein, 5 (v/v)% fetal calf serum, at 37 °C for 4 days on a cover glass placed in a Petri dish, in the same manner as in Reference Example 4 which will be described later. With respect to the protein produced by the cell culture of each of the above-obtained clones, determination of NANBV antigenicity was conducted by indirect fluorescent antibody technique using specific antisera described in Step 2. As a result, it was found that the protein produced in the G418 resistant cell clone reacted with all of the antisera of guinea pigs immunized with the NANBV-related antigens obtained in Reference Example 1 (see Table 6).

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Table 6

Detection by fluorescent antibody technique of NANBV-related antigens produced in transformant human or chimpanzee hepatocyte

		from 1	NANB atient	guinea pig antise		
cell extract	pooled serum	acute	chronic	anti- core	anti- NS3	anti- NS5
human hepatocy	yte-der	ived				
HL-Al	+	+	+	+	+	+
HL-A2	+	. +	+	+	+	+
normal Chang Liver	-	-	-	_	••	-
chimpanzee he	patocyt	e-deri	ved			
CL-B11	+	+	+	+	+	+
CL-B14	+	+	+	+	+	+
normal Chimp Liver	_	-	-	-	-	-

This fact means that the entire coding region of NANBV gene covering the region coding for the core antigen through the region coding for NS5 was expressed.

Step 4 (Sucrose density-gradient centrifugation of NANBV-related antigens produced by human hepatocyteand chimpanzee hepatocyte-derived transformant cell clones HL-A1 and CL-B11)

Clones HL-A1 and CL-B11 were individually cultured t 37 °C for 4 days on 5 Petri dishes having a diameter of 9 cm in a  $\rm CO_2$  incubator, in the same manner as in Step 3. The cells were scraped off with a rubber policeman and pooled together with the culture liquid and subjected to ultrasonic treatment at 20 kHz (200 W) for 2 minutes and centrifugation at 5000 x g and at 4 °C for 15 minutes. The resultant supernatant was further subjected to centrifugation at 48000 x g and at 4 °C for 14 hours to obtain a precipitate. The precipitate was suspended in 1 ml of M/75 PBS and subjected to ultrasonic treatment for 2 minutes and sucrose density-gradient centrifugation at 160000 x g and at 4 °C for 15 hours, followed by fractionation. Each fraction was subjected to SDS-polyacrylamide electrophoresis and Western blotting analysis in the same manner as in Step 2, thereby conducting the detection of a core antigen and an envelope antigen. As a result, both antigens were detected at a sucrose density of 40 (w/v)% to 50 (w/v)%, indicating that there were obtained NANBV particles.

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#### Example 2

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Step 1 (Construction of a plasmid for the expression in yeast of the entire coding region of the NANBV genomic cDNA and preparation of a transformant yeast)

Plasmid pORF24 obtained in Step 1 of Example 1 (Part 11) was digested with Xbal to thereby obtain an NANBV cDNA fragment of about 9 kb in the same manner as in Step 1 of Example 1 (Part II). 0.5 μg of this cDNA fragment was dissolved in a T4 DNA polymerase solution containing 67 mM Tris-HCl (pH 8.8), 6.7 mM magnesium chloride, 16.6 mM ammonium sulfate, 10 mM 2-ME, 6.7 mM EDTA-2Na 0.02 (w/v)% bovine albumine and 0.3 mM dNTP, and 2 to 5 units of T4 DNA polymerase (Takara Shuzo Co., Ltd., Japan) was added thereto and the resultant mixture was incubated at 37 °C for 60 minutes, thereby rendering blunt the both terminals of the fragment. Then, Xhol linker (CCTCGAGG) was ligated thereto by means of T4 DNA ligase. Illustratively stated, 0.3 µg of the DNA was dissolved in 21 µl of a 10 x ligation solution containing 500 mM Tris-HCl (pH 7.5), 100 mM magnesium chloride, 100 mM DTT and 10 mM ATP. Added to the resultant mixture were 300 to 400 units of T4 DNA ligase (Takara Shuzo Co., Ltd., Japan) and distilled water in an amount such that the total volume became 210  $\mu$ l, followed by incubation at 14 °C for 18 hours. The thus prepared cDNA fragment was inserted in an expression vector for use in yeast, namely, YEp133PCT (described in U.S. Patent No. 4,810,492) at its Xhol site to thereby obtain expression plasmid pYHC5. With this expression plasmid pYHC5, yeast S. cerevisiae (ATCC No. 44772) was transformed by the alkali cation method (Ito, H. et al, J. Bacteriol., 153:163-168, 1983), to thereby obtain transformant yeast YHC5-1. This transformant was designed so that when the culture medium was lack of phosphate ions, the gene for repressive acid phosphatase was activated to cause the transcription of the NANBV cDNA ligated downstream thereof, thereby producing NANBV-related antigens.

Step 2 (production of NANBV-related antigens by yeast and characterization thereof)

Transformant yeast YHC5-1 obtained in Step 1 was inoculated into 100 ml of a culture medium prepared by adding 20 µg/ml of each of uracil, L-tryptophan and L-histidine into Burkholder's medium (see Burkholder, P.R. et al, Am.J. Botany, 30, 206-211, 1943) that was a totally synthesized medium containing 1.5 g/l of potassium phosphate monobasic. The inoculated medium was cultured at 30 °C for 24 hours while shaking. The cultured yeast was washed with physiological saline and inoculated into 1000 ml of a fresh medium of the same type as described above except that 1.5 g/l of potassium chloride was contained instead of potassium phosphate monobasic. The inoculated medium was cultured at 30 °C for 24 hours and the resultant cells were collected. The collected cells were suspended in M/75PBS and glass beads (diameter: 0.45-0.55 mm) were added thereto and the suspension was subjected to a Bead Beater (manufactured and sold by Biospec Products, U.S.A.) to thereby disrupt the cells, and then to centrifugation at 10000 x g and at 4 °C for 10 minutes, thereby obtaining a supernatant. The thus obtained supernatant was subjected to SDS-polyacrylamide gel electrophoresis and Western blotting analysis in the same manner as in Step 2 of Example 1 (Part II), thereby examining whether or not NANBV-related antigens had been produced. As a result, it was found that the extract of transformant yeast YHC5-1 reacted with all of the antibodies respectively specific for the core antigen, envelope antigen, NS3 protein and NS5 protein. This fact means that the entire cording region of NANBV gene from the core antigen region through the NS5 region was expressed (see Table 7).

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# Table 7 Reactivity of proteins produced by transformant yeast YHC5-1 with NANBV-related antibodies

		serum hepat:	guinea pig antiser				
10	cell extract	pooled	acute	chronic	anti- core	anti- NS3	anti- NS5
15	YHC5-1	+*	+	+	+	+	+
	normal S.cerevisiae	-	•	•	-	-	-

 Reactivity as measured by Western blotting analysis

Further, the cell extract was subjected to sucrose density-gradient centrifugation in the same manner as in Step 4 of Example 1 (Part II). As a result, both of the core antigen and the envelope antigen were detected in a fraction at a sucrose density of 40 (w/v)% to 50 (w/v)%, indicating that there were obtained NANBV particles.

#### 30 Example 3

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Step 1 (Construction of a plasmid for introduction into vaccinia virus)

Plasmid pUV1 (Falko G. Falkner, Sekhar Chakrabarti and Bernard Moss; Nucleic Acid Res., 15 (17), 7192, 1987) was digested with restriction enzyme EcoRI and subjected to phenol extraction and ethanol precipitation, thereby obtaining a DNA. In the same manner as in Reference Example 1, 0.5 μg of this DNA was dissolved in a T4 DNA polymerase solution and 2 to 5 units of T4 DNA polymerase (manufactured and sold by Takara Shuzo Co., Ltd., Japan) was added, followed by incubation at 37 °C for 60 minutes, to thereby render blunt both terminals. Separately, a DNA fragment carrying the entire nucleotide sequence of NANBV gene coding for an NANBV protein was obtained by a method in which plasmid pORF-24 described in Step 1 of Example 1 (Part II) was digested with Xbal or with Xbal and EcoRI to thereby obtain a DNA fragment of about 9 kb or a DNA fragment of 6.4 kb and the obtained DNA fragment was then treated with T4 DNA polymerase, thereby rendering both terminals blunt, 0.5 µg of the thus obtained DNA derived from pUV1 and 0.5 µg of the thus obtained DNA derived from pORF-24 were dissolved in 21 µl of 10 x ligation solution in the same manner as in Reference Example 1 and added thereto were 300 to 400 units of T4 DNA ligase (manufactured and sold by Takara Shuzo Ltd., Japan) and distilled water in an amount such that the total volume became 210 µl, followed by incubation at 14 °C for 12 to 18 hours. Thus, a cDNA derived from NANBV was ligated to pUV1 at its EcoRI site located downstream of the promoter. The ligation reaction mixture was subjected to phenol extraction and the aqueous layer was subjected to ethanol precipitation to collect a DNA. With the DNA, E. coli strain JM109 was transformed in accordance with the calcium chloride method, as described in Step 6 of Example 1 (Part I), thereby obtaining plasmid clones pXX-49 and pXX-51 each having an NANBV cDNA fragment of about 9 kb. In addition, plasmid pXE-39 having an NANBV cDNA fragment of about 6.4 kb and lacking the NS5 region of NANBV. The results are shown in Fig. 5.

# 55 Reference Example 4

(Culturing of vaccinia virus WR strain)

Vaccinia virus WR strain was cultured by the customary method. Illustratively stated, monolayer-cultured

cells [such as mouse-derived thymidine kinase (TK)-defective cell line L-M(TK<sup>-</sup>) (ATCC CCL-1.3, Dainippon Pharmaceutical Co., Ltd., Japan), simian kidney-derived Vero cells and adult human hepatocyte Chang Liver (ATCC CCL-13, Dainippon Pharmaceutical Co., Ltd., Japan)] were cultured in a Petri dish having a diameter of 6 cm. The resultant cells were inoculated with 0.5 ml of vaccinia virus and allowed to stand at 37 °C for 1 to 2 hours, followed by removing the virus liquid. Then, 5 ml of Eagle's MEM medium (manufactured and sold by Nissui Pharmaceutical Co., Ltd., Japan) having, added thereto, 5 (v/v)% fatal calf serum and the cells were cultured at 37 °C for 24 to 48 hours until satisfactory cytopathiceffect was observed. Then, the virus culture liquid or the infected cells were collected and suspended in MEM and subjected to ultrasonic treatment at 20 kHz (200W) for 2 minutes, thereby obtaining a virus liquid.

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#### Refernce Example 5

(Infectivity assay for vaccinia virus by plaque method)

The plaque method was conducted by the customary procedure. Illustratively stated, the cell culture described in Reference Example 4 in a 6 cm-diameter Petri dish is inoculated with the virus liquid obtained in Reference Example 4, which had been diluted 10-fold with M-199 (manufactured and sold by Sigma, U.S.A.), in an amount of 0.1 ml/dish and allowed to stand at 37 °C for 2 hours, thereby adsorbing the virus onto the cells. Then, the inoculum liquid was removed and an agar-containing medium (prepared by adding 3 (v/v)% fetal calf serum, 0.14 (w/v)% NaHCO<sub>3</sub> and 0.8 (w/v)% agar to M-199) was added in an amount of 5 ml/dish. After the agar had solidified at room temperature, culturing was conducted at 37 °C for 24 hours. Then a medium prepared by adding neutral red (manufactured and sold by Wako Pure Chemical Industries, Ltd., Japan) to the above-mentioned agar in an amount of about 0.006 (w/v)% was overlaid in an amount of 2.5 ml/dish, followed by further culturing at 37 °C. The number of the resultant plaques was counted, to thereby determine the infectivity.

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#### Step 2 (Preparation of recombinant vaccinia virus)

Vero cells were cultured for 24 hours in a 9 cm-dlameter Petri dish to be used for tissue culture (manufactured and sold by Falcon, U.S.A.). The cultured Vero cells were inoculated with vaccinia virus WR strain (ATCC VR-119) at an MOI (multiplicity of infection) of 0.05 and allowed to stand at 37 °C for 2 hours so as to adsorb the virus onto the cells. After that period, the virus liquid was removed and the cells were washed with MEM twice. Then, 1 µg and 5 µg of the plasmid DNAs obtained in Step 1 of Example 3 were individually subjected to calcium phosphate precipitation of DNA in accordance with the calcium phosphate method (Graham, F.L., van der Eb, A.J.: Virology, 52, 456-467, 1973), thereby obtaining 1 ml of a DNA-calcium phosphate precipitation solution with respect to each plasmid DNA. The thus obtained 1 ml each of DNA-calcium phosphate precipitate solution was added to the above-obtained virus-infected cells and allowed to stand at room temperature for 30 minutes. Then, 15 ml of the virus culture medium as described in Reference Example 4 was added and the resultant mixture was incubated at 37 °C for 3.5 hours. Then the virus culture medium was removed and 15 mt of a fresh virus culture medium was added, followed by culturing at 37 °C for 48 hours. Then the cell culture was subjected to freezing and thawing 3 times, to thereby obtain a virus suspension. The virus suspension was inoculated into L-M(TK<sup>-</sup>) cells cultured in a 6 cm-diameter Petri dish to be used for culturing, in the same manner as in Reference Example 5. After the virus had been adsorbed onto the cells, an agar medium containing 25 μg/ml of 5-Bromo-2'-deoxyuridine (BUdR, manufactured and sold by Sigma, U.S.A.) was added in an amount of 5 ml/dish, followed by culturing at 37 °C for 8 hours. Then, an agar medium containing 25 μg/ml of BUdR and 25 μg/ml of 5-Bromo-4-Chloro-3-Indolyl-β-D-Galactoside (X-Gai, Takara Shuzo Co., Ltd., Japan) was overlald in an amount 2.5 ml/dlsh and the cells were further cultured at 37 °C for 2 days. The blue plaques that appeared were collected together with the overlaying agar medium and suspended in 0.5 ml of M-199 and the supermatant was inoculated into L-M(TK') cells in the same manner as described above, followed by plaque doning 3 times to thereby purify the clones. The thus obtained vaccinia virus was a recombinant vaccinia virus transformed by recombination with a plasmid vector DNA having an NANBV cDNA, and had the β-galactosidase gene and lacked thymidine kinase. The results are shown in Table 8.

Table 8
Characteristics of recombinant vaccinia virus

recombinant	employed	NANBV genome	thymidine kinase activity	β-galacto sidase activity	
virus clone	combination	(kb)	activity	accivicy	
vUV17	pUV1	x	x	•0	
vUV27	pUV1	x	x	0	
VXE17	pXD39	0(6.4)	x	0	
VXE28	pXE39	0(6.4)	X	0	
vXX19	pXX49	0(9.0)	x	0	
vXX29	pXX51	0(9.0)	x	0	
vxx39	pXX51	0(9.0)	x	0	

Note) O: Observed

X : Not observed

35 Step 3 (Detection and confirmation by fluorescent antibody technique of production of NANBV-related antigens by recombinant vaccinia virus)

Antigens produced by recombinant vaccinia virus were detected and confirmed by indirect fluorescent antibody technique. L-M(TK1) cells were cultured on a cover glass and inoculated with recombinant vaccinia virus which had been diluted 10-fold, and the cells were cultured by the method described in Reference Example 4. After culturing for 48 hours, the cover glass was taken out and washed with M/75 phosphate buffer saline (M/75 PBS) (pH 7.4) three times and then with distilled water one time, followed by air-drying, Then, the cells were fixed by acetone at -20 °C for 5 minutes. An anti-NANBV mouse monoclonal antibody to be used as a primary antibody was obtained from a hybridoma obtained by fusing a mouse myeloma cell with a lymphocyte separated from BALD/C mouse immunized by the customary method using the core antigen and non-structural protein antigens NS-3 and NS-5. A monoclonal antibody specific for the envelope was prepared by using an antigen produced by binding a 16-mer oligopeptide to bovine serum albumin, which 16-mer oligopeptide was comprised of an amino acid sequence (1st to 16th amino acids on the N-terminal side) deduced from the envelope gene. Indirect fluorescent antibody technique was conducted by the customary method. That is, the infected cells fixed by acetone were reacted with the primary antibody at 37 °C for 1 hour and washed with M/75 PBS three times. Then, the cells were reacted with FITC (fluorescent dye)-labeled anti-human or mouse IgG antibody (manufactured and sold by Cappel Co., Ltd., Germany) at 37 °C for 1 hour and washed with M/75 PBS three times, followed by the observation by a fluorescence microscope. The results are shown in Table 9.

Table 9 (1)

Detection by fluorescent antibody technique of
NANBV-related antigens produced by cells infected
with recombinant vaccinia virus

10		serum from healthy human			serum from NANB hepatitis patient					
15	recombinant vaccinia virus	#6	#8	#9	pooled serum #II-1	pooled serum #PS-1	acute	chronic		
20	vUV17	-	•	-	-	-	÷	-		
20	v <b>u</b> V27	-	-	-	-	-	-	-		
25	vXE17	-	-	_	+	+	+	+		
	vXE28	-	-	-	+	+	+	+		
30	<b>vXX19</b> .	_	_	_	+	+	+	+		
	<b>v</b> XX29	-	_	-	+	· +	+	+		
35	vXX39	-	-	-	+	+	+	+		

(to be continued)

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Table 9 (2) (continued)

5	recombinant				
	vaccinia virus	anti-core #11	anti-EnV #755	anti-NS3 #74-1	anti-NS5 #8905
10	vUV17	**	•	-	•
	vUV27	· 🛥	-	-	-
15		•			
	VXE17	+	+	+ ,	-
	vXE28	+	+	+	. <del>-</del>
20				•	
	vXX19	+	+	+	+
	vXX29	+	+	+	+
25	vXX39	+	+	<b>+</b>	+

Recombinant vaccinia virus clones vXE17 and vXE28, both of which lacked a portion of the nucleotide sequence coding for the NS5 region of NANBV, and clones vXX19, vXX29 and vXX39, all of which had a complete ORF coding for the protein of NANBV, reacted with all of the sera from NANB hepatitis patients but did not react with any of the sera from healthy humans. When a mouse monoclonal antibody was employed, all of clones vXE17 and vXE28 and clones vXX19, vXX29 and vXX39 reacted with the anti-core antigen monoclonal antibody, the anti-envelope antigen monoclonal antibody and the anti-NS3 monoclonal antibody. With respect to the anti-NS5 monoclonal antibody, vXE17 and vXE28, both of which lacked the NS5 region of NANBV, did not reacted therewith, but vXX19, vXX29 and vXX39 reacted therewith. These facts mean that the desired expression products were advantageously produced by means of recombinant vaccinia virus and that particularly by means of clones vXX19, vXX29 and vXX39, the entire region was expressed, from the core antigen on the N-terminal side of the NANBV protein through the NS5 protein on the C-terminal side thereof.

Step 4 (Analysis of supernatant of culture of cells infected with recombinant vaccinia virus by sucrose density-gradient centrifugation)

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As described in Reference Example 4, 1.0 ml (1.2 x 10<sup>7</sup> PFU) of recombinant vaccinia virus vXX39 (1.2 x 10<sup>7</sup> PFU/ml) was inoculated into a cell culture of human hepatocyte Chang Liver and adsorbed onto the cells at 37 °C for 2 hours and then cultured at 37 °C for 3 days with only M-199. 200 ml of the supernatant of the culture was subjected to centrifugation at 3000 x g for 5 minutes to obtain a supernatant, which was further subjected to centrifugation at 48000 x g and at 4 °C for 14 hours, to obtain a precipitate. The precipitate was suspended in 2 ml of M/75 PBS and subjected to ultrasonic treatment at 20 kHz (200W) for 2 minutes and the resultant product was subjected to sucrose density-gradient centrifugation at 160,000 x g and at 4 °C for 15 hours and wherein the sucrose density was changed from 20 (w/v)% to 60 (w/v)%, thereby obtaining fractions. Each of the fractions was mixed with a sample buffer containing, in the final concentration, 20 (v/v)% glycerol, 100 mM (pH 6.8) Tris HCl, 2 (w/v)% SDS, 2 (v/v)% 2-mercaptoethanol and 0.02 (w/v)% BPB, and heated at 100 °C for 5 minutes and then subjected to 0.1 (w/v)% SDS-12.5 % polyacrylamide gel electrophoresis, to thereby separate proteins from each other. Then the gel was subjected to a trans blot apparatus (manufactured and sold by Nippon Eido Co., Ltd., Japan), to thereby blot the proteins which were separated by electrophoresis onto Hybond-ECL membrane (manufactured and sold by Amershan, England). The Hybond-ECL membrane

was then immersed in a solution composed of 10 mM Tris HCl (pH 7.2), 150 mM NaCl, 0.05 (w/v)% Tween-20 (T-TBS) and 5 (w/v)% sklm milk, and incubated at room temperature for 1 hour, thereby blocking the membrane. Then, the anti-core antigen monoclonal antibody (clone 29) which had been diluted 500-fold with a T-TBS buffer containing 1 % skim milk was reacted with the Hybond-ECL membrane at 37 °C for 1 hour and then the membrane was washed well 2 times with a fresh one of the above-mentioned T-TBS buffer containing 1 % skim milk. Then, the membrane was reacted with biotin-labeled anti-mouse IgG (manufactured and sold by Cappel Co., Itd., Germany; diluted 500-fo1d) at room temperature for 1 hour. The Hybond-ECL membrane was then washed 2 times and reacted with HRPO-labeled streptoavidin (manufactured and sold by Amersham, England; diluted 500-fold) at room temperature for 1 hour, and washed well 4 times with T-TBS. The membrane was subjected to chemical luminescene reaction by means of ECL Western blotting detection system (manufactured and sold by Amersham, England). The membrane was wrapped with Saran Wrap and kept in contact with an X-ray film for 30 seconds, followed by the development of the film. As a result, it was found that activities of a core antigen and an envelope antigen of NANBV were observed at a sucrose concentration of 44 to 58 (w/v)% (as shown in Fig. 7). This fact indicates that there were obtained NANBV particles.

Step 5 (Observation of recombinant vaccinia virus-infected cells under electron microscope)

Recombinant vaccinia virus clones vXX39 and vUV17 (the later having no NANBV genome) were cultured on human hepatocyte for 2 days in the same manner as described in Reference Example 4, and the infected cells were collected by means of a rubber police. The collected cells were embedded in an epoxy resin (manufactured and sold by Nissin EM Co., Ltd., Japan) by the customary method and ultra-thin section samples were prepared by slicing. The samples were subjected to uranium-lead double staining, using 2 % uranium acetate and lead citrate, and examined under an electron microscope. As a result, particles were observed in the cells infected with vXX39 as shown in Fig. 8, but not in the cells infected with vUV17 and having no NANBV genome in the cytoplasm. These results, taken together with the results of Step 4, show that vXX39 had produced NANBV particles.

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5 SEQUENCE LISTING 10 -(1) GENERAL INFORMATION: (1) APPLICANT: THE RESEARCH FOUNDATION OF MICROBIAL DISEASES OF OSAKA UNIVERSITY 15 (11) TITLE OF INVENTION: NON-A, NON-B HEPATITIS VIRUS GENOMIC CONA AND ANTIGEN POLYPEPTIDE (iii) NUMBER OF SEQUENCES: 2 (iv) CORRESPONDENCE ADDRESS: 20 (A) ADDRESSEE: Brookes & Martin (B) STREET: High Holborn House, 52/54, High Holborn (C) CITY: London (E) COUNTRY: United Kingdom (F) ZIP: WC1V 6SE 25 (v) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 30 (vi) CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: EP 91305717.0 (B) FILING DATE: 25-JUNE-1991 (C) CLASSIFICATION: (vii) PRIOR APPLICATION DATA: 35 (A) APPLICATION NUMBER: JP 2-167466 (B) FILING DATE: 25-JUN-1990 (vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: JP 2-230921 (B) FILING DATE: 31-AUG-1990 (vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: JP 2-305605 (B) FILING DATE: 09-NOV-1990 45 (vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: US 07/635,451 (B) FILING DATE: 28-DEC-1990 (vii) PRIOR APPLICATION DATA:

(A) APPLICATION NUMBER: JP 3-132090

(B) FILING DATE: 08-MAY-1991

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(vii) PRIOR APPLICATION DATA: 10 (A) APPLICATION NUMBER: JP 3-138493 (B) FILING DATE: 14-MAY-1991 (viii) ATTORNEY/AGENT INFORMATION: (A) NAME: BLAKE, John H. (C) REFERENCE/DOCKET NUMBER: JHB/91-1011 15 (1x) TELECOMMUNICATION INFORMATION: (A) TELEPHONE: (44) 892-510600 (B) TELEFAX: (44) 71-831-0586 20 (2) INFORMATION FOR SEQ ID NO:1: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 9416 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double 25 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (genomic) (111) HYPOTHETICAL: NO 30 (iv) ANTI-SENSE: NO (v) FRAGMENT TYPE: internal (vi) ORIGINAL SOURCE: (A) ORGANISM: Hepatitis virus 35 (B) STRAIN: Non-A, Non-B (C) INDIVIDUAL ISOLATE: Human (D) DEVELOPMENTAL STAGE: Suspension cells (E) HAPLOTYPE: Diploid (F) TISSUE TYPE: Liver (G) CELL TYPE: Hepatocyte 40 (H) CELL LINE: ATCC CCL 13 (vii) IMMEDIATE SOURCE: (A) LIBRARY: BK170, BK171, BK172 (B) CLONE: pDEL-NS5 45 (viii) POSITION IN GENOME: (A) CHROMOSOME/SEGMENT: N/A (B) MAP POSITION: Infectious Agent (C) UNITS: bp (ix) FEATURE: 50 (A) NAME/KEY: CDS

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## (B) LOCATION: 333..9362

	(	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	:1;							
	CGAT	TGGG	GG C	GACA	CT.CC	A CC	ATAG.	ATCA	стс	CCCT	GTG A	AGGA.	ACTA	CT G	TCTT	CACGC		60
15	AGAA	AGCG	TC T.	AGCC	ATGG	C GT	TAGT.	ATGA	GTG	TCGT	GCA 1	GCCT	CCAG	GA C	cccc	ССТСС		120
	CGGG	AGAG	CC A	TAGT	GGTC	T GC	GGAA	CCGG	TGA	GTAC	ACC	GGAA	TTGC	CA G	GACG	ACCGG		180
	GTCC	TTTC	TT G	GATC	AACC	c gc	TCAA	TGCC	TGG	AGAT	TTG	GGCG	TGCC	cc c	GCGA	GACTG		240
20	CTAG	CCGA	GT A	GTGT	TGGG	T CG	CGAA	AGGC	CTT	GTGG	TAC	TGCC	TGAT	AG G	GTGC	TTGCG		300
	AGTG	cccc	GG G	AGGT	CTCG	T AG	ACCG	TGCA	CC	ATG Met 1	AGC Ser	ACG Thr	AAT Asn	CCT . Pro 5	AAA Lys	CCT Pro		353
25	CAA	AGA	AAA	ACC	AAA	CGT	AAC	ACC	AAC	CGC	CGC	CCA	CAG	GAC	GTC	AAG		401
	Gln	Arg	Lys 10	Thr	Lys	Arg	Asn	Thr 15	Asn	Arg	Arg	Pro	G1n 20	Asp	Val	Lys		
30	TTC Phe	CCG	GGC	GGT	GGT	CAG	ATC	GTT	GGT	GGA	GTT	TAC	CTG	TTG	CCG	CGC		449
	Pne	25	ч	GFY	GIY	GIŅ	30	vai	uly	Giy	141	35	ren	Leu	110	מיש		
	AGG	GGC	ccc	AGG	TTG	GGT	GTG	CGC	GCG	ccc	AGG	AAG	ACT	TCC	GAG	CGG		497
35	Arg 40	Gly	Pro	Arg	Leu	45	VAI	AFG	AIA	Pro	50	Lys	1111	361	414	55		
	TCG	CAA	CCT	CGT	GGA	AGG	CGA	CAA	CCT	ATC	CCC	AAG	GCT	CGC	CGG	CCC	٠,	545
	Ser	GIN	Pro	Arg	60 60	Arg	Arg	GIN	PFO	65	Pro	Lys	Ala	VI A	70	rio	, ,	
40	GAG	GGC	AGG	ACC	TGG	GCT	CAG	CCC	GGG	TAC	CCT	TGG	CCT	CTC	TAT	GGC		593
	Glu	GIY	Arg	Thr 75	∤rp	Ala	GIN	Pro	80	ıyr	PTO	קזו	FIU	85	131	diy		
	AAT	GAG	GGC	TTA	GGG	TGG	GCA	GGA	TGG	СТС	CTG	TCA	CCC	CGC	GGC	TCC		641
45	ASN	GIU	90 90	Leu	GIY	irp	Ala	95		Lea	reu	261	100	Alg	uly	361		
	CGG	ССТ	AGT	TGG	GGC	CCC	ACG	GAC	CCC	CGG	CGT	AGG	TCG	CGT	AAT	TTG		689
	Arg	Pro 105		Trp	GIY	Pro	110		Pro	AFG	arg	115		MIÐ	noil	Lou		
50	GGT	AAG	GTC	ATC	GAT	ACC	СТС	ACA	TGC	GGC	TTC	GCC	GAT	CTC	ATG	GGG		737
	Gly	Lys	val	TIE	ASP	וחר	reu	100	cys	uly	rne	AIB	42h	rea	ne t	Gly		

TAC ATT CCG CTC GTC GGC GCC CCC CTG GGG GGC GCT GCC AGG GCC CTG Tyr Ile Pro Leu Val Gly Ala Pro Leu Gly Gly Ala Ala Arg Ala Leu GCA CAT GGT GTC CGG GTT CTG GAG GAC GGC GTG AAC TAT GCA ACA GGG Ala His Gly Val Arg Val Leu Glu Asp Gly Val Asn Tyr Ala Thr Gly AAT CTG CCC GGT TGC TCT TTT TCT. ATC TTC CTC TTG GCT CTG CTG TCC Asn Leu Pro Gly Cys Ser Phe Ser Ile Phe Leu Leu Ala Leu Leu Ser TGC CTG ACC ACC CCA GCT TCC GCT TAC GAA GTG CAC AAC GTG TCC GGG Cys Leu Thr Thr Pro Ala Ser Ala Tyr Glu Val His Asn Val Ser Gly ATA TAT CAT GTC ACG AAC GAC TGC TCC AAC GCA AGC ATT GTG TAT GAG Ile Tyr His Val Thr Asn Asp Cys Ser Asn Ala Ser Ile Val Tyr Glu GCA GCG GAC TTG ATC ATG CAT ACT CCT GGG TGC GTG CCC TGC GTT CGG Ala Ala Asp Leu Ile Met His Thr Pro Gly Cys Val Pro Cys Val Arg GAA GGC AAC TCC TCC CGC TGC TGG GTA GCG CTC ACT CCC ACG CTC GCA Glu Gly Asn Ser Ser Arg Cys Trp Val Ala Leu Thr Pro Thr Leu Ala GCC AGG AAC GTC ACC ATC CCC ACC ACG ACG ATA CGA CGC CAC GTC GAT Ala Arg Asn Val Thr Ile Pro Thr Thr Thr Ile Arg Arg His Val Asp CTG CTC GTT GGG GCG GCT GCT TTC TGT TCC GCT ATG TAC GTG GGG GAC Leu Leu Val Gly Ala Ala Ala Phe Cys Ser Ala Met Tyr Val Gly Asp CTC TGC GGA TCT GTT TTC CTC GTC TCT CAG CTG TTC ACC TTC TCG CCT Leu Cys Gly Ser Val Phe Leu Val Ser Gln Leu Phe Thr Phe Ser Pro CGC CGG CAT GTG ACA TTA CAG GAC TGT AAC TGC TCA ATT TAT CCC GGC Arg Arg His Val Thr Leu Gln Asp Cys Asn Cys Ser Ile Tyr Pro Gly CAT GTG TCG GGT CAC CGT ATG GCT TGG GAC ATG ATG AAC TGG TCG His Val Ser Gly His Arg Met Ala Trp Asp Met Met Asn Trp Ser 

10	9					CTA Leu												1361
4	5					GTG Val												1409
,	•					ATG Met												1457
2	0	CTA Leu	CTT	77T Phe	GCT Ala	GGC Gly 380	GTT Val	GAÇ Asp	GGG Gly	GAT Asp	ACC Thr 385	CAC His	GTG Val	ACA Thr	GGG Gly	GGG G1y 390	GCG Ala	1505
•	æ					ACC Thr												1553
2	5					CAG G1n												1601
3	10	AGG Arg	ACT Thr 425	GCC Ala	CTG Leu	AAC Asn	TGC Cys	AAT Asn 430	GAC Asp	TCT Ser	CTC Leu	CAG Gln	ACT Thr 435	GGG Gly	TTT Phe	CTT Leu	GCC Ala	1649
	05	GCG Ala 440	CTG Leu	TTC Phe	TAC Tyr	ACA Thr	CAT His 445	AGT Ser	TTC Phe	AAC Asn	TCG Ser	TCC Ser 450	GGG Gly	TGC Cys	CCA Pro	GAG G1u	CGC Arg 455	1697
•	· .	ATG Met	GCC Ala	CAG Gln	TGC Cys	CGC Arg 460	ACC Thr	ATT Ile	GAC Asp	AAG Lys	TTC Phe 465	GAC Asp	CAG Gln	GGA Gly	TGG Trp	GGT Gly 470	CCC Pro	1745
	10	ATT	ACT Thr	TAT Tyr	GCT Ala 475	GAG Glu	TCT Ser	AGC Ser	AGA Arg	TCA Ser 480	Asp	CAG Gln	AGG Arg	CCA Pro	TAT Tyr 485	TGC Cys	TGG Trp	1793
	45	CAC His	TAC Tyr	CCA Pro 490	Pro	CCA Pro	CAA Gln	TGT Cys	ACC Thr 495	Ile	GTA Val	CCT Pro	GCG Ala	TCG Ser 500	Glu	GTG Val	TGC Cys	1841
•	<del>,,</del> ,	GGC Gly	CCA Pro 505	Val	TAC	TGC Cys	TTC Phe	ACC Thr 510	Pro	AGC Ser	CCT Pro	GTC Val	GTC Val 515	۷al	GGG Gly	ACG Thr	ACC Thr	1889
	50	GAT Asp	CGT	TTC Phe	GGT	GTC Val	CCT	ACG Thr	TAT Tyr	AGA Arg	TGG Trp	GGG Gly	GAG Glu	AAC Asn	GAG Glu	ACT Thr	GAC Asp	1937

40	520					525	•				530					535	
10	GTG Val	CTG Leu	CTG Leu	CTC Leu	AAC Asn 540	AAC Asn	ACG Thr	CGG Arg	CCG Pro	CCG Pro 545	CAA G1n	GGC Gly	AAC Asn	TGG Trp	TTC Phe 550	GGC Gly	1985
15	TGC Cys	ACA Thr	TGG Trp	ATG Met 555	AAT Asn	AGC Ser	ACC Thr	GGG Gly	TTC Phe 560	ACC Thr	AAG Lys	ACA Thr	TGT Cys	GGG G1y 565	GGG Gly	CCC Pro	2033
20	CCG Pro	TGT Cys	AAC Asn 570	ATC Ile	GGG G1y	GGG Gly	GTC Val	GGC Gly 575	AAC Asn	AAC Asn	ACC Thr	CTG Leu	ACC Thr 580	TGC Cys	ccc Pro	ACG Thr	2081
20	GAC Asp	TGC Cys 585	TTC Phe	CGG Arg	AAG Lys	CAC His	CCC Pro 590	GAG G1u	GCT Ala	ACC Thr	TAC Tyr	ACA Thr 595	AAA Lys	TGT Cys	GGT Gly	TCG Ser	2129
25	GGG Gly 600	Pro	TGG Trp	CTG Leu	ACA Thr	CCT Pro 605	AGG Arg	TGC Cys	ATG Met	GTT Val	GAC ASP 61D	TAT Tyr	CCA Pro	TAC Tyr	AGG	CTC Leu 615	2177
30	TGG Trp	CAT His	TAC Tyr	CCC Pro	TGC Cys 620	Thr	GTT Val	AAC Asn	TTT Phe	ACC Thr 625	He	TTC Phe	AAG Lys	GTT Val	AGG Arg 630	Met	2225
30	TAT Tyr	GTG Val	GGG Gly	GGG Gly 635	GTG Val	GAG Glu	CAC His	AGG Arg	CTC Leu 640	Asn	GCT A1a	GCA Ala	TGC Cys	AAT Asn 645	Trp	ACC Thr	2273
35	ÇGA Arg	GGA Gly	GAG Glu 650	Arg	TGT Cys	GAC Asp	TTG Leu	GAG G1u 655	GAC Asp	AGG Arg	GAT Asp	AGG Arg	CCG Pro 660	Glu	CTC Leu	AGC Ser	2321
40	CCG Pro	CTG Leu 665	Leu	CTG Leu	TCT Ser	ACA Thr	ACA Thr 670	G1 u	TG0	CAG Gln	GTA Val	CTG Leu 675	Pro	TGT	TCC Ser	TTC Phe	2369
40	ACC Thr 680	Thr	CTA Leu	CCA Pro	GCT Ala	CTG Leu 685	Ser	ACT Thr	GGC Gly	teu	ATT Ile 690	His	CTC Leu	CAT His	CAG Glr	AAC Asn 695	2417
45	ATC Ile	GTC Val	GAC Asp	GTG Val	CAA Glr 700	Tyr	CTA Let	TAC Tyr	GG1 G1	ATA 11e 705	Gly	TCA Ser	GCG Ala	GTT Val	GT0 Val 710	C TCC Ser	2465
50	TT1 Phe	GCA Ala	ATC a Ile	AAA B Lys 715	Trp	GAG Glu	TAT I Tyr	gT(	C CT( 1 Let 72(	Lei	CTT Lei	TTC Phe	CTT	CT( Let 725	, Let	A GCG u Ala	2513

1 <b>0</b>		GCA Ala															2561
15		GAG Glu 745	Ala														2609
		GGC Gly															2657
20		TAC Tyr															2705
25		GTG Val															2753
*		GCC Ala															2801
30		GGT Gly 825															2849
35	GCT Ala 840	AGG Arg	CTC Leu	ATA Ile	TGG Trp	TGG Trp 845	TTA Leu	CA'A G1n	TAT Tyr	TTT Phe	ACC Thr 850	ACC Thr	AGA Arg	GCC Ala	GAG Glu	GCG Ala 855	2897
<b>30</b>		TTA Leu									Ala						2945
40		ATC								His							2993
	ATC Ile	ACC Thr	AAA Lys 890	Leu	CTA Leu	ATT I le	GCC Ala	ATA Ile 895	CTC Leu	GGT Gly	CCG Pro	CTC	ATG Met 900	GTG Val	CTC	CAA Gìn	3041
45	GCT Ala	GGC Gly 905	Ile	ACC Thr	AGA Arg	GTG Val	CCG Pro 910	TAC Tyr	TTC Phe	GTG Val	CGC Arg	GCT Ala 915	GIn	GGG Gly	CTC Leu	ATT Ile	3089
50	CAT H1s	GCA Ala	TGC Cys	ATG Met	TTA	GTG Val	CGG Arg	AAG Lys	GTC Val	GCT	GGG Gly	GGT Gly	CAT	TAT Tyr	GTC Val	CAA Gln	3137

			*		
10	920	925		930	935
				GGC ACG TAC ATT TAC Gly Thr Tyr Ile Tyr 950	
15				GCG GGC CTA CGA GAC Ala Gly Leu Arg Asp 965	
20		. Val Glu Pro V		GAC ATG GAG ACC AAG ASP Met Glu Thr Lys 980	
		Gly Ala Asp 1		GGG GAC ATC ATC TTG Gly Asp Ile Ile Leu 995	
25	CTG CCC GTC Leu Pro Val 1000	TCC GCC CGA / Ser Ala Arg / 1005	Arg Gly Lys Glu	ATA CTC CTG GGC CCG Ile Leu Leu Gly Pro 1010	GCC 3377 Ala 1015
30				CTC GCG CCC ATC ACG Leu Ala Pro Ile Thr 103	Ala
	TAC TCC CAA Tyr Ser Glr	CAG ACG CGG ( Gin Thr Arg ( 1035	GGC CTA CTT GGT Gly Lau Lau Gly 1040	TGC ATC ATC ACT AGC Cys lle lle Thr Ser 1045	CTT 3473 Leu
35	ACA GGC CGC Thr Gly Arg 105	g Asp Lys Asn (	CAG GTC GAG GGA Gln Val Glu Gly 1055	GAG GTT CAG GTG GTT Glu Val Gln Val Val 1060	TCC 3521 Ser
40	ACC GCA ACA Thr Ala Thi 1085	r Gln Ser Phe	CTG GCG ACC TGC Leu Ala Thr Cys 1070	GTC AAC GGC GTG TGT Val Asn Gly Val Cys 1075	TGG 3569 Trp
	ACC GTT TAG Thr Val Tyr 1080	C CAT GGT GCT : r His Gly Ala : 1085	Gly Ser Lys Thr	TTA GCC GCG CCA AAG Leu Ala Ala Pro Lys 1090	GGG 3617 Gly 1095
45	CCA ATC ACI Pro Ile Th	C CAG ATG TAC r Gln Met Tyr 1100	ACT AAT GTG GAC Thr Asn Val Asp 1105	CAG GAC CTC GTC GGC Gln Asp Leu Val Gly 111	Trp
	CCC AAG CCC Pro Lys Pro	C CCC GGG GCG D Pro Gly Ala 1115	CGT TCC TTG ACA Arg Ser Leu Thr 1120	CCA TGC ACC TGT GGC Pro Cys Thr Cys Gly 1125	AGC 3713 Ser
60					

10	Ser Asp L	TT TAC TTG eu Tyr Leu 130	Val Thr Ar	GA CAT GCT : rg His Ala :	GAC GTC ATT Asp Val Ile 1140	Pro Val Arg	3761
15	CGG CGG G Arg Arg G 1145	GC GAC AGT Ty Asp Ser	AGG GGG AG Arg Gly Se 1150	GC CTG CTC er Leu Leu	TCC CCC AGG Ser Pro Arg 1155	CCT GTC TCC Pro Val Ser	3809
	TAC TTG A Tyr Leu L 1160	ys Gly Ser	TCG GGT GG Ser Gly Gl 1185	ly Pro Leu	CTC TGC CCC Leu Cys Pro 1170	TTC GGG CAC Phe Gly His 1175	3857
20	GCT GTG G Ala Val G	GC ATC TTC Bly Ile Phe 1180	Arg Ala Al	CC GTA TGC la Val Cys 1185	ACC CGG GGG Thr Arg Gly	GTT GCG AAG Val Ala Lys 1190	3905
25					GAA ACT ACT Glu Thr Thr		3953
	Pro Val P	TTC ACG GAC Phe Thr Asp 1210	Asn Ser Se	CC CCC CCG er Pro Pro 215	GCC GTA CCG Ala Val Pro 1220	Gln Ser Phe	4001
30	CAA GTG G Gln Val A 1225	GCC CAC CTA Ala His Leu	CAC GCT CC His Ala Pr 1230	CC ACT GGC ro Thr Gly	AGC GGC AAG Ser Gly Lys 1235	AGT ACT AAA Ser Thr Lys	4049
35					AAG GTG CTC Lys Val Leu 1250		4097
33	CCG TCC C Pro Ser \	GTT GCC GCT Val Ala Ala 1260	Thr Leu G	GG TTT GGG ly Phe Gly 1265	Ala Tyr Met	TCT AAG GCA Ser Lys Ala 1270	4145
40					GTA AGG ACC Val Arg Thr		4193
	Gly Ala i	CCC GTC ACA Pro Val Thr 1290	Tyr Ser T	CC TAT GGC hr Tyr Gly 295	AAG TTT CTT Lys Phe Leu 1300	GCC GAT GGT Ala Asp Gly )	4241
45	GGT TGC 1 Gly Cys 5 1305	Ser Gly Gly	GCT TAT G Ala Tyr A 1310	AC ATC ATA	ATA TGT GAT Ile Cys Asp 1315	GAG TGC CAT Glu Cys H1s	4289
50	TCA ACT (	GAC TCG ACT Asp Ser Thr	ACA ATC T Thr Ile L	TG GGC ATC eu Gly Ile	GGC ACA GTC Gly Thr Val	CTG GAC CAA Leu Asp Gln	4337

	1320	1325 ·	1330	1335
10	GCG GAG ACG GCT GGA Ala Glu Thr Ala Gly 134	Ala Arg Leu Val Val	G CTC GCC ACC GCT ACG I Leu Ala Thr Ala Thr 15 1350	Pro
15	CCG GGA TCG GTC ACC Pro Gly Ser Val Thr 1355	QTG CCA CAC CCA AAC Val Pro His Pro Asr 1360	C ATC GAG GAG GTG GCC n lle Glu Glu Val Ala 1365	CTG 4433 Leu
	TCT AAT ACT GGA GAG Ser Asn Thr Gly Glu 1370	ATC CCC TTC TAT GGC   Ile Pro Phe Tyr Gly   1375	C AAA GCC ATC CCC ATT y Lys Ala Ile Pro Ile 1380	GAA 4481 Glu
20	GCC ATC AGG GGG GGA Ala Ile Arg Gly Gly 1385	AGG CAT CTC ATT TTC Arg His Leu Ile Pho 1390	C TGT CAT TCC AAG AAG e Cys His Ser Lys Lys 1395	AAG 4529 Lys
25	TGC GAC GAG CTC GCC Cys Asp Glu Leu Ala 1400	GCA AAG CTG TCA GG Ala Lys Leu Ser Gl 1405	C CTC GGA ATC AAC GCT y Leu Gly Ile Asn Ala 1410	GTG 4577 Val 1415
	GCG TAT TAC CGG GGG Ala Tyr Tyr Arg Gly 142	/ Leu Asp Val Ser Va	C ATA CCA ACT ATC GGA l lle Pro Thr lle Gly 25 143	Asp
30	GTC GTT GTC GTG GCA Val Val Val Val Ala 1435	A ACA GAC GCT CTG AT Thr Asp Ala Leu Me 1440	G ACG GGC TAT ACG GGC t Thr Gly Tyr Thr Gly 1445	GAC 4673 Asp
35	TTT GAC TCA GTG ATC Phe Asp Ser Val Ile 1450	GAC TGT AAC ACA TG ASP Cys Asn Thr Cy 1455	T GTC ACC CAG ACA GTC s Val Thr Gln Thr Val 1460	GAC 4721 Asp
	TTC AGC TTG GAT CCC Phe Ser Leu Asp Pro 1465	C ACC TTC ACC ATT GA Thr Phe Thr Ile Gl 1470	G ACG ACG ACC GTG CCT u Thr Thr Val Pro 1475	CAA 4769 Gln
40	GAC GCA GTG TCG CGG Asp Ala Val Ser Arg 1480	C TCG CAG CGG CGG GG g Ser Gln Arg Arg Gl 1485	T AGG ACT GGC AGG GGT Y Arg Thr Gly Arg Gly 1490	AGG 4817 Arg 1495
45	AGA GGC ATC TAC AGA Arg Gly Ile Tyr Arg 150	g Phe Val Thr Pro Gl	AA GAA CGQ CCC TCG GGC ly Glu Arg Pro Ser Gly 505 15	/ Met
50	TTC GAT TCC TCG GT Phe Asp Ser Ser Va 1515	C CTG TGT GAG TGC TA 1 Leu Cys Glu Cys Ty 1520	AT GAC GCG GGC TGT GCT Vr Asp Ala Gly Cys Ala 1525	TTGG 4913 a Trp
<b>∞</b>			7	

		,		
10	TAC GAG CTC ACC CCG GC Tyr Glu Leu Thr Pro Al 1530	C GAG ACC TCG GTT a Glu Thr Ser Val 1535	AGG TTG CGG GCC TAC CTG Arg Leu Arg Ala Tyr Leu 1540	4961
	AAC ACA CCA GGG TTG CC Asn Thr Pro Gly Leu Pr 1545	C GTT TGC CAG GAC o Val Cys Gln Asp 1550	CAC CTG GAG TTC TGG GAG His Leu Glu Phe Trp Glu 1555	5009
15	Ser Val Phe Thr Gly Le	C ACC CAT ATA GAT u Thr His Ile Asc 65	GCA CAC TTC TTG TCC CAG Ala His Phe Leu Ser Gln 1570 1575	5057
20	ACC AAG CAG GCA GGA GA Thr Lys Gln Ala Gly At 1580	C AAC TTC CCC TAC Sp Asn Phe Pro Typ 158	C CTG GTA GCA TAC CAA GCC r Leu Val Ala Tyr Gln Ala 35 1590	5105
	ACG GTG TGC GCC AGG G Thr Val Cys Ala Arg A 1595	CT CAG GCC CCA CC la Gln Ala Pro Pro 1600	T CCA TCA TGG GAT CAA ATG D Pro Ser Trp Asp Gln Met 1605	5153
25	TGG AAG TGT CTC ATA C Trp Lys Cys Leu Ile A 1610	GG CTG AAA CCT AC rg Leu Lys Pro Th 1615	G CTG CAC GGG CCA ACA CCC r Leu His Gly Pro Thr Pro 1620	5201
30	TTG CTG TAC AGG CTG G Leu Leu Tyr Arg Leu G 1625	GA GCC GTC CAG AA ly Ala Val Gln As 1630	T GAG GTC ACC CTC ACC CAC n Glu Val Thr Leu Thr His 1635	5249
35	Pro Ile Thr Lys Tyr I	TC ATG GCA TGC AT le Met Ala Cys Me 645	G TCG GCT GAC CTG GAG GTC it Ser Ala Asp Leu Glu Val 1650 1655	5297
	GTC ACT AGC ACC TGG C Val Thr Ser Thr Trp V 1660	al Leu Val Gly Gl	A GTC CTT GCA GCT CTG GCC y Val Leu Ala Ala Leu Ala 65 1670	5345
40	GCG TAT TGC CTG ACA A Ala Tyr Cys Leu Thr 1 1675	CA GGC AGT GTG GT hr Gly Ser Val Va 1680	TC ATT GTG GGT AGG ATT ATC al lie Val Gly Arg lie lie 1685	5393
45	TTG TCC GGG AGG CCG ( Leu Ser Gly Arg Pro / 1690	CC ATT GTT CCC G/ la Ile Val Pro As 1695	AC AGG GAG CTT CTC TAC CAG SP Arg Glu Leu Leu Tyr Gln 1700	5441
	GAG TTC GAT GAA ATG ( Glu Phe Asp Glu Met ( 1705	GAA GAG TGC GCC TO Glu Glu Cys Ala So 1710	CG CAC CTC CCT TAC ATC GAG er His Leu Pro Tyr Ile Glu 1715	5489
50	CAG GGA ATG CAG CTC	GCC GAG CAA TTC A	AG CAG AAA GCG CTC GGG TTA ys Gln Lys Ala Leu Gly Leu	5537

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	1720	1725	1730	1735
10	CTG CAA ACA GCC Leu Gln Thr Ala	ACC AAA CAA GO Thr Lys Gln A 1740	CG GAG GCT GCT C la Glu Ala Ala Ala F 1745	ccc gTG GTG GAG 5585 Pro Val Val Glu 1750
15	TCC AAG TGG CGA Ser Lys Trp Arg 175	Ala Leu Glu Ti	CA TTC TGG GCG AAG ( hr Phe Trp Ala Lys F 1760	CAC ATG TGG AAT 5633 His Met Trp Asn 1765
	TTC ATC AGC GGG Phe Ile Ser Gly 1770	Ile Gln Tyr L	TA GCA GGC TTA TCC / eu ala Gly Leu Ser 775	ACT CTG CCT GGG 5681 Thr Leu Pro Gly 1780
20	AAC CCC GCA ATA Asn Pro Ala Ile 1785	GCA TCA TTG A Ala Ser Leu M 1790	TG GCA TTC ACA GCC let Ala Phe Thr Ala 1795	TCT ATC ACC AGC 5729 Ser Ile Thr Ser
25	CCG CTC ACC ACC Pro Leu Thr Thr 1800	CAA AGT ACC C Gln Ser Thr L 1805	CTC CTG TTT AAC ATC Leu Leu Phe Asn Ile 1810	TTG GGG GGG TGG 5777 Leu Gly Gly Trp 1815
	GTG GCT GCC CAA Val Ala Ala Glr	CTC GCC CCC C Leu Ala Pro F 1820	CCC AGC GCC GCT TCG Pro Ser Ala Ala Ser 1825	GCT TTC GTG GGC 5825 Ala Phe Val Gly 1830
30	GCC GGC ATC GCC Ala Gly Ile Ala 183	a Gly Ala Ala '	GTT GGC AGC ATA GGC Val Gly Ser Ile Gly 1840	CTT GGG AAG GTG 5873 Leu Gly Lys Val 1845
35	CTT GTG GAC AT Leu Val Asp Ile 1850	e Leu Ala Gly	TAT GGA GCA GGA GTG Tyr Gly Ala Gly Val 1855	GCC GGC GCG CTC 5921 Ala Gly Ala Leu 1850
40	GTG GCC TTT AA Val Ala Phe Ly: 1865	G GTC ATG AGC s Val Met Ser 1870	GGC GAG ATG CCC TCC Gly Glu Met Pro Ser 187	Thr Glu ASP Leu
	GTC AAT CTA CT Val Asn Leu Le 1880	T CCT GCC ATC u Pro Ala Ile 1885	CTC TCT CCT GGC GCC Leu Ser Pro Gly Ala 1890	CTG GTC GTC GGG 6017 Leu Val Val Gly 1895
45	GTC GTG TGT GC Val Val Cys Al	A GCA ATA CTG a Ala Ile Leu 1900	CGT CGA CAC GTG GGT Arg Arg His Val Gly 1905	CCG GGA GAG GGG 6065 Pro Gly Glu Gly 1910
50	Ala Val Gln Tr	G ATG AAC CGG p Met Asn Arg	CTG ATA GCG TTC GCC Leu Ile Ala Phe Ala 1920	TCG CGG GGT AAT 6113 Ser Arg Gly Asn 1925

10	CAT GTT His Val	TCC CCC Ser Pro 1930	ACG CAC	TAT GTO Tyr Vai 193	Pro G	AG AGC G Tu Ser A	AC GCC G sp Ala A 1940	CA GCG la Ala		6161
	GTT ACT Val Thr 194	Gln Ile	CTC TCC Leu Ser	AGC CTT Ser Let 1950	ACC AT	TC ACT C le Thr G 1	AG CTG C In Leu Li 955	TG AAA eu Lys		6209
15	CTC CAC Leu His 1960	CAG TGG Gln Trp	ATT AAT Ile Asn 196	Glu Ası	C TGC TO Cys So	CC ACA C er Thr P 1970	CG TGT TO	er Gly	TCG Ser 1975	6257
20	TGG CTA Trp Leu	AGG GAT Arg Asp	GTT TGG Val Trp 1980	GAC TG	o Ila C	GC ACG G ys Thr V 985	TG TTG A	CT GAC hr Asp 1990	Phe	6305
25	AAG ACC Lys Thr	TGG CTC Trp Leu 199	Gin Ser	AAG CT	C CTG C u Leu P 2000	CG CAG C	eu Pro G	GA GTC ly Val 005	CCT P <b>ro</b>	6353
25	TTT TTC Phe Phe	TCG TGC Ser Cys 2010	CAA CGC Gln Arg	GGG TA Gly Ty 20	r Lys G	GA GTC T	GG CGG G rp Arg G 2020	GA GAC	GGC Gly	6401
30	ATC ATG Ile Met 202	Gin Thr	ACC TGC	CCA TG Pro Cy 2030	T GGA G s Gly A	ICA CAG A la Gln I	TC ACC G le Thr G	GA CAT	GTC Val	6449
35	AAA AAC Lys Asn 2040	GGT TCC Gly Ser	ATG AGG Met Arg 204	Ile Va	C GGG C	CCT AAG A Pro Lys 1 2050	ACC TGC A Thr Cys S	GC AAC Ger Asn	ACG Thr 2055	6497
	TGG CAT	GGA ACA	TTC CCC Phe Pro 2060	ATC AAD Ile As	n Ala I	TAC ACC A Tyr Thr 1 2065	ACG GGC C Thr Gly F	CCC TGC Pro Cys 2070	Thr	6545
40	CCC TCT Pro Ser	CCA GCG Pro Ala 207	Pro Asi	TAT TO 1 Tyr Se	T AGG ( r Arg A 2080	GCG CTG	Trp Arg \	STG GCC /al Ala 2085	GCT Ala	6593
45	GAG GAG Glu Glu	TAC GTO Tyr Val 2090	GAG GTO	1 Thr Ar	G GTG 0 g Val 0 195	GGG GAT G	TTC CAC 1 Phe His 1 2100	rac GTG Tyr Val	ACG Thr	6641
	GGC ATO	: The Thi	GAC AA Asp As	C GTA AA n Val Ly 2110	G TGC ( 's Cys I	CCA TGC ( Pro Cys (	CAG GTT ( Gln Val F 2115	CCG GCT Pro Ala	CCT Pro	6689
50	GAA TTO	TTC TCC	GAG GT	G GAC GC 1 Asp G	A GTG (	CGG TTG Arg Leu	CAC AGG THIS Arg	TAC GCT Tyr Ala	CCG Pro	6737

GCG TGC AGG CCT CTC CTA CGG GAG GAG GTT ACA TTC CAG GTC GGG CTC Ala Cys Arg Pro Leu Leu Arg Glu Glu Val Thr Phe Gln Val Gly Leu AAC CAA TAC CTG GTT GGG TCA CAG CTA CCA TGC GAG CCC GAA CCG GAT Asn Gin Tyr Leu Vai Gly Ser Gin Leu Pro Cys Glu Pro Glu Pro Asp GTA GCA GTG CTC ACT TCC ATG CTC ACC GAC CCC TCC CAC ATC ACA GCA Val Ala Val Leu Thr Ser Met Leu Thr Asp Pro Ser His Ile Thr Ala GAA ACG GCT AAG CGT AGG TTG GCC AGG GGG TCT CCC CCC TCC TTG GCC Glu Thr Ala Lys Arg Arg Leu Ala Arg Gly Ser Pro Pro Ser Leu Ala AGC TCT TCA GCT AGC CAG TTG TCT GCG CCT TCC TTG AAG GCG ACA TGC Ser Ser Ser Ala Ser Gln Leu Ser Ala Pro Ser Leu Lys Ala Thr Cys ACT ACC CAC CAT GTC TCT CCG GAC GCT GAC CTC ATC GAG GCC AAC CTC Thr Thr His His Val Ser Pro Asp Ala Asp Leu Ile Glu Ala Asn Leu CTG TGG CGG CAG GAG ATG GGC GGG AAC ATC ACC CGC GTG GAG TCG GAG Leu Trp Arg Gln Glu Met Gly Gly Asn Ile Thr Arg Val Glu Ser Glu AAC AAG GTG GTA GTC CTG GAC TCT TTC GAC CCG CTT CGA GCG GAG GAG Asn Lys Val Val Val Leu Asp Ser Phe Asp Pro Leu Arg Ala Glu Glu GAT GAG AGG GAA GTA TCC GTT CCG GCG GAG ATC CTG CGG AAA TCC AAG Asp Glu Arg Glu Val Ser Val Pro Ala Glu Ile Leu Arg Lys Ser Lys AAG TTC CCC GCA GCG ATG CCC ATC TGG GCG CGC CCG GAT TAC AAC CCT Lys Phe Pro Ala Ala Met Pro Ile Trp Ala Arg Pro Asp Tyr Asn Pro CCA CTG TTA GAG TCC TGG AAG GAC CCG GAC TAC GTC CCT CCG GTG GTG Pro Leu Leu Glu Ser Trp Lys Asp Pro Asp Tyr Val Pro Pro Val Val CAC GGG TGC CCG TTG CCA CCT ATC AAG GCC CCT CCA ATA CCA CCT CCA His Gly Cys Pro Leu Pro Pro Ile Lys Ala Pro Pro Ile Pro Pro Pro

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10	Arg Arg				Thr Glu	TCC TCC ( Ser Ser )			7361
15		Glu Leu				AGC TCC ( Ser Ser ( 2355			7409
,,	_			Thr Ala		GAC CAG ( Asp Gln / 2370			7457
20				-		TCC TCC / Ser Ser i	Met Pro		7505
25			Gly Asp			GAC GGG '		Ser Thr	7553
20					Val Val	TGC TGC			7601
30		Thr Gly				GCT GCG ( Ala Ala ( 2435			7649
35				Ser Asn		CTG CGC ( Leu Arg ( 2450			7697
-						CTG CGG ( Leu Arg (			7745
40			Leu Gln			CAC TAC		Val Leu	7793
45					Thr Val	AAG GCT . Lys Ala			7841
		Glu Ala				CAT TCG His Ser 2515	Ala Lys		7889
50						CTA TCC			7937

2525 · AAC CAC ATC CAC TCC GTG TGG AAG GAC TTG CTG GAA GAC ACT GTG ACA Asn His Ile His Ser Val Trp Lys Asp Leu Glu Asp Thr Val Thr CCA ATT GAC ACC ACC ATC ATG GCA AAA AAT GAG GTT TTC TGT GTC CAA Pro Ile Asp Thr Thr Ile Met Ala Lys Asn Glu Val Phe Cys Val Gln CCA GAG AAA GGA GGC CGT AAG CCA GCC CGC CTT ATC GTA TTC CCA GAT Pro Glu Lys Gly Gly Arg Lys Pro Ala Arg Leu Ile Val Phe Pro Asp CTG GGA GTC CGT GTA TGC GAG AAG ATG GCC CTC TAT GAT GTG GTC TCC Leu Gly Val Arg Val Cys Glu Lys Met Ala Leu Tyr Asp Val Val Ser ACC CTT CCT CAG GTC GTG ATG GGC TCC TCA TAC GGA TTC CAG TAC TCT Thr Leu Pro Gln Val Val Met Gly Ser Ser Tyr Gly Phe Gln Tyr Ser CCT GGG CAG CGA GTC GAG TTC CTG GTG AAT ACC TGG AAA TCA AAG AAA Pro Gly Gln Arg Val Glu Phe Leu Val Asn Thr Trp Lys Ser Lys Lys AAC CCC ATG GGC TTT TCA TAT GAC ACT CGC TGT TTC GAC TCA ACG GTC Asn Pro Met Gly Phe Ser Tyr Asp Thr Arg Cys Phe Asp Ser Thr Val ACC GAG AAC GAC ATC CGT GTT GAG GAG TCA ATT TAC CAA TGT TGT GAC Thr Glu Asn Asp Ile Arg Val Glu Glu Ser Ile Tyr Gln Cys Cys Asp TTG GCC CCC GAA GCC AGA CAG GCC ATA AAA TCG CTC ACA GAG CGG CTT Leu Ala Pro Glu Ala Arg Gln Ala Ile Lys Ser Leu Thr Glu Arg Leu TAT ATC GGG GGT CCT CTG ACT AAT TCA AAA GGG CAG AAC TGC GGT TAT Tyr Ile Gly Gly Pro Leu Thr Asn Ser Lys Gly Gln Asn Cys Gly Tyr CGC CGG TGC CGC GCG AGC GGC GTG CTG ACG ACT AGC TGC GGT AAC ACC Arg Arg Cys Arg Ala Ser Gly Val Leu Thr Thr Ser Cys Gly Asn Thr CTC ACA TGT TAC TTG AAG GCC TCT GCA GCC TGT CGA GCT GCG AAG CTC Leu Thr Cys Tyr Leu Lys Ala Ser Ala Ala Cys Arg Ala Ala Lys Leu

10	GAC Asp		Thr					Gly					Va1			8561
	AGC Ser 2745	Ala					Asp					Arg				8609
15	GCT Ala					Ser					Asp					8657
20	TAC Tyr				Leu					Ser					Val	8705
25	CAC His			Ser					Tyr					Asp		8753
25	ACC Thr		Leu					Trp					His			8801
30	AAC Asn 282	Ser					Ile					Pro				8849
35	AGG Arg D					Thr					Ile					8897
	CAA G1n				Ala					Ile					Tyr	8945
40	ATT Ile			Leu					Ile					His		8993
45	AGC Ser		Phe					Tyr					Ile			9041
	GCT Ala 290	Ser					Leu					Leu				9089
50	CAT H1s															9137

	2920	2925	2930	2935
10	AGG GCC GCC ACT TGT Arg Ala Ala Thr Cys 294	GGC AAA TAC CTC TTC GGly Lys Tyr Leu Phe 10 2945	Asn Trp Ala Val Lys	Thr
15	AAA CTT AAA CTC ACT Lys Leu Lys Leu Thr 2955	CCA ATC CCG GCT GCG Pro Ile Pro Ala Ala 2960	TCC CGG CTG GAC TTG Ser Arg Leu Asp Leu 2965	TCC 9233 Ser
20	GGC TGG TTC GTT GCT Gly Trp Phe Val Ala 2970	GGT TAC AGC GGG GGA Gly Tyr Ser Gly Gly 2975	GAC ATA TAT CAC AGC Asp Ile Tyr His Ser 2980	CTG 9281 Leu
20	TCT CGT GCC CGA CCC Ser Arg Ala Arg Pro 2985	C CGT TGG TTC ATG CTG Arg Trp Phe Met Leu 2990	TGC CTA CTC CTA CTT Cys Leu Leu Leu Leu 2995	TCT 9329 Ser
25	GTA GGG GTA GGC ATO Val Gly Val Gly Ilo 3000	C TAC CTG CTC CCC AAC e Tyr Leu Leu Pro Asn 3005	CGA TGAACGGGGA GATA Arg 3010	AACACT 9382
	CCAGGCCAAT AGGCCAT	ccc cetttititt titt		9416
30	(2) INFORMATION FO	R SEQ ID NO:2:		
35	(A) L (B) T	E CHARACTERISTICS: ENGTH: 3010 amino aci YPE: amino acid OPOLOGY: linear	ds	
	(ii) MOLECUL	E TYPE: protein		
	(x1) SEQUENC	E DESCRIPTION: SEQ ID	NO:2:	
40	Met Ser Thr Asn Pr	o Lys Pro Gln Arg Lys 5 10		Asn.
	Arg Arg Pro Gln As	p Val Lys Phe Pro Gly 25	Gly Gly Gln Ile Val	Gly
45	Gly Val Tyr Leu Le 35	eu Pro Arg Arg Gly Pro 40	o Arg Leu Gly Val Arg 45	, Ala
	Pro Arg Lys Thr Se 50	er Glu Arg Ser Gln Pro 55	o Arg Gly Arg Arg Gli 60	n Pro
50	Ile Pro Lys Ala Ar 65	rg Arg Pro Glu Gly Arg	g Thr Trp Ala Gln Pro 75	5 Gly 80

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10	Tyr	Pro	Trp	Pro	Leu 85	Tyr	G·l y	Asn	G1u	G1y 90	Leu	Gly	Trp.	Ala	G1y 95	Trp	
	Leu	Leu	Ser	Pro 100	Arg	Gly	Ser		Pro 105	Ser	Trp	Gly	Pro	Thr 110	Asp	Pro	
15	Arg	Arg	Arg 115	Ser	Arg	Asn	Len	G1y 120	Lys	۷a٦	Ile	Asp	Thr 125	Leu	Thr	Cys	
	Gly	Phe 130	Ala	Asp	Leu	Met	Gly 135	Tyr	Ide	Pro	Leu	Va1 140	Gly	Ala	Pro	Leu	
20	Gly 145	Gly	Ala	Ala	Arg	A1a 150	Leu	Æla	His	Gly	Val 155	Arg	Val	Leu	Glu	ASP 180	
	Gly	Val	Asn	Tyr	A1a 165	Thr	G1y	Asn	Leu	Pro 170	Gly	Cys	Ser	Phe	Ser 175	I Je	
25	Phe	Leu	Leu	Ala 180	Leu	Leu	Ser	Cys	Leu 185	Thr	Thr	Pro	Ala	Ser 190	Ala	Туг	
	Glu	Val	His 195	Asn	Va1	Ser	Gly	Ile 200	Tyr	His	Val	Thr	Asn 205	Asp	Cys	Ser	
30	Asn	Ala 210		Ile	Val	Tyr	Glu 215		Ala	Asp	Leu	11e 220	Met	His	Thr	Pro	
	Gly 225	Cys	Val	Pro	Суз	Va1 230	Arg	Glu	GTy	Asn	Ser 235	Ser	Arg	Cys	Trp	Val 240	
35	Ala	Leu	Thr	Pro	Thr 245	Leu	Ala	Ala	Arg	Asn 250	Val	Thr	. IJe	Pro	Thr 255	Thr	
	Thr	· Ile	Arg	Arg 260		Val	Asp	Leu	Leu 265	Val	'G1y	Ala	Ala	A1a 270	Phe	Cys	
40	Ser	- Ala	Met 275		Val	Gly	Asp	Leu 280		Gly	Ser	Va1	Phe 285	Leu	Va1	Ser	
	Gìr	1 Leu 290		Thr	Phe	Ser	Pro 295		Arg	His	≀ Va¹	Thr 300	· Leu	G1n	Asp	Cys	
45	AS:		s Ser	Ile	Туг	9rc 310		/ His	va1	Ser	- Gly 31!	/ His S	s Arg	) Met	. Ala	320	
	Ası	o Met	t Met	. Met	325		Şe:	- Pro	Thr	7 Thi 330	r Ala	a Lei	Va¹	Val	335	- Gln	
50	Le	u Lei	u Arg	340 340		G1r	A18	a Va	1 Va 34	) Asp	Me1	t Va	1 Ala	350	/ Ala	His	

10	Trp	Gly	Va 1 355	Leu	Ala	Gly	Leu	A1a 360	Туг	Tyr	Ser	Met	Ala 365	Gly	Asn	Trp	
	Ala	Lys 370	۷a۱	Leu	Ile	Val	Met 375	Leu	Leu	Phe	Ala	G1y 380	Val	Asp	Gly	Asp	
15	Thr 385	His	۷a٦	Thr	Gly	Gly 390	Ala	Gln	Ala	Lys	Thr 395	Thr	Asn	Arg	Leu	Va 1 400	
	Ser	Met	Phe	Ala	Ser 405	Gly	Pro	Ser	Gln	Lys 410	Ile	G1n	Leu	Ile	Asn 415	Thr	
20	Asn	Gly	Ser	420	His		Asn	Arg	Thr 425	Ala	Leu	Asn	Çys	Asn 430	Asp	Ser	
	Leu	Gln	Thr 435	G1y	Phe	Leu	Ala	A1a 440	Leu	Phe	Tyr	Thr	H1s 445	Ser	Phe	Asn	
25	Ser	Ser 450	Gly	Cys	Pro	Glu	Arg 455	Met	Ala	Gln	Cys	Arg 460	Thr	Ile	Asp	Lys	
	Phe 465	Asp	Gln	G1y	Trp	Gly 470		Ile	Thr	Tyr	A1a 475		Ser	Ser	Arg	Ser 480	
30	Asp	Gln	Arg	Pro	Tyr 485	Cys	Trp	His	Туг	Pro 490	Pro	Pro	G1n	Cys	Thr 495	Ile	
	Val	Pro	Ala	Ser 500		Val	Cys	Gly	Pro 505	۷a۱	Tyr	Cys	Phe	Thr 510	Pro	Ser	
35	Pro	Val	Va 1 515		Gly	Thr	Thr	Asp 520		Phe	Gly	Va]	Pro 525	Thr	Туг	Arg	
	Trp	Gly 530		Asn	Glu	Thr	Asp 535	Val	Leu	Leu	Leu	Asn 540	Asn	Thr	Arg	Pro	
40	Pro 545		1, G1y	' Asn	Trp	9 Phe 550		Cys	Thr	Trp	Met 555	Asn	Ser	Thr	Gly	Phe 560	
	Thr	Lys	Thr	· Cys	Gly 565		Pro	) Pro	Cys	Asr 570	ı Ile	G1y	Gly	/ Val	G1y 575	Asn	
45	Asr	Thi	Let	1 Thr 580		Pro	Thr	. Asp	Cys 585		e Arg	Lys	His	590	G1u	Ala	
	Thr	Туі	r Thr 595		s Cys	G1)	/ Ser	Gly 600		Tr	Leu	Thr	Pro 605	Arg	Cys	Met	
50	۷a	Ası 61		- Pro	у Туі	- Arg	5 Lei 61		His	: Ту	r Pro	620	Thi	r Val	Asn	Phe	

	Thr 625	Ile	Phe	Lys	Val	Arg 630	Met	Tyr	Va-1	Gly	G1y 635	Va1	Glu	His	Arg	Leu 640
10	Asn	Ala	Ala	Cys	Asn 645	Trp	Thr	Arg	Gly	G1บ 650	Arg	Cys	Asp	Leu	Glu 655	Åsp
15	Arg	Asp	Arg	Pro 660	Glu	Leu	Ser	Pro	Leu 665	Leu	Leu	Ser	Thr	Thr 670	Glu	Trp
,,,	G1n	Val	Leu 675	Pro	Cys	Ser	Phe	Thr 680	Thr	Leu	Pro	Ala	Leu 685	Ser	Thr	Gly
20	Leu	Ile 690	His	Leu	His	Gln	Asn 695	Ile	Val	Asp	Val	G1n 700	Tyr	Leu	Tyr	Gly
	Ile 705	Gly	Ser	Ala	Val	Val 710	Ser	Phe	Ala	Ile	Lys 715	Trp	Glu	Tyr	Val	Leu 720
25	Leu	Leu	Phe	Leu	Leu 725	Leu	Ala	Asp	Ala	Arg 730	Val	Cys	Ala	Cys	Leu 735	Trp
	Met	Met	Leu	Leu 740	Ile	Ala	Gln	Ala	G1u 745	Ala	Ala	Leu	Glu	Asn 750	Leu	Val
30	Val	Leu	Asn 755	Ser	Ala	Ser	Val	Ala 760	Gly	Ala	His	Gly	Ile 765	Leu	Ser	Phe
	Leu	Va1 770	Phe	Phe	Cys	Ala	Ala 775	Trp	Tyr	Ile	Lys	Gly 780	Arg	Leu	Val	Pro
35	Gly 785	Ala	Thr	Tyr	Ala	Leu 790	Tyr	Gly	Val	Trp	Pro 795	Leu	Leu	Leu	Leu	Leu 800
	Leu	Ala	Leu	Pro	Pro 805	Arg	Ala	Tyr	Ala	Met 810	Asp	Arg	Glu	Met	A1a 815	Ala
40	Ser	Cys	Gly	Gly 820	Ala	Val	Phe	Val	G1y 825	Leu	Val	Leu	Leu	Thr 830	Leu	Ser
	Pro	Tyr	Tyr 835	Lys	Val	Phe	Leu	A1a 840	Arg	Leu	Ile	Trp	Trp 845	Leu	Gln	Туг
45	Phe	Thr 850	Thr	Arg	Ala	Glu	A1a 855	Asp	Leu	His	Val	Trp 860	Ile	Pro	Pro	Leu
	Asn 865		Arg	Gly	Gly	Arg 870		Ala	Ile	Ile	Leu 875	Leu	Met	Cys	Ala	Val 880
50	His	Pro	Glu	Leu	Ile 885	Phe	Asp	Ile	Thr	Lys 890		Leu	Ile	Ala	Ile 895	Leu

10	Gly	Pro	Leu	Met 900	Val	Leu	G1n	Ala	Gly 905	Ile	Thr	Arg	Val	Pro 910	Туг	Phe
	Val	Arg	Ala 915	Gln	Gly	Leu	Ile	His 920	Ala	Cys	Met	Leu	Va1 925	Arg	Lys	Va1
15	A1a	Gly 930	Gly	His	Tyr	Val	G1n 935	Met	Ala	Phe	Met	Lys 940	Leu	Gly	Ala	Leu
	Thr 945	Gly	Thr	Tyr	lle	Tyr 950	Asn	His	Leu	Thr	Pro 955	Leu	Arg	Asp	Trp	Pro 960
20	Arg	Ala	Gly	Leu	Arg 965	Asp	Leu	Ała	Val	Ala 970	Val	Glu	Pro	Val	Val 975	Phe
	Ser	Asp	Met	G1u 980	Thr	Lys	Ile	Ile	Thr 985	Trp	Gly	Ala	Asp	Thr 990	Ala	Ala
25	Cys	Gly	Asp 995	Ile	Ile	Leu	Gly	Leu 1000		Val	Ser	Ala	Arg 100	_	Gly	Lys
	Glu	Ile 1010		Leu	Gly	Pro	Ala 1015		Ser	Leu	Glu	Gly 1020		Gly	Leu	Arg
30	Leu 1025	Leu	Ala	Pro	Пe	Thr 1030		Tyr	Ser	G1n	G}n 1035		Arg	Gly	Leu	Leu 1040
	Gly	Cys	Ile	Ile	Thr 1045		Leu	Thr	Gly	Arg 105(		Lys	Asn	Gln	Va1 105	
35	Gly	Glu	Va1	G1n 1060		Val	Ser	Thr	Ala 106		G1n	Ser	Phe	Leu 1076		Thr
	Cys	Va1	Asn 1075		Va1	Cys	Trp	Thr 1080		Tyr	His	Gly	A1a 108		Ser	Lys
40	Thr	Leu 1090		Ala	Pro	Lys	Gly 109		He	Thr	Gln	Met 1100		Thr	Asn	Val
	Asp 110	Gln 5	Asp	Leu	Val	G1y 1110		Pro	Lys	Pro	Pro 111		Ala	Arg	Ser	Leu 1120
<b>4</b> 5	Thr	Pro	Cys	Thr	Cys 112		Ser	Ser	Asp	Leu 1130		Leu	Val	Thr	Arg 113	
	Ala	Asp	Val	I le 1140		Val	Arg	Arg	Arg 114		Asp	Ser	Arg	Gly 115	Şer O	Leu
50	Leu	Ser	Pro 115		Pro	Val	Ser	Tyr 116		Lys	Gly	Ser	Ser 116		G1y	Pro

10	leu	Leu 1170		Pro	Phe	Gly	His 1175	A'la	Val	Gly	Ile	Phe 1180	_	Ala	Ala	Val
	Cys 1185		Arg	Gly	Val	Ala 1190	•	Ala	Va <sup>-</sup> 1	Asp	Phe 1195		Pro	Val	Glu	Ser 1200
15	Met	Glu	Thr	Thr	Met 1205		Sér	Pro	Val	Phe 1210		Asp	Asn	Ser	Ser 1215	
	Pro	Ala	Val	Pro 1220		Ser	Phe	Głn	Val 1225		His	Leu	His	Ala 1230		Thr
20	Gly	Ser	Gly 1235		Ser	Thr	Lys	Va 1 1240		Ala	Ala	Tyr	A1a 1245		Gln	Gly
	Tyr	Lys 1250		Leu	Val	Leu	Asn 1255	Pro	Ser	Val	Ala	Ala 1260		Leu	Gly	Phe
25	Gly 1265		Tyr	Met	Ser	Lys 1270		His	Gĩy	Ile	Asp 1275		Asn	Ile	Arg	Thr 1280
	Gly	Val	Arg	Thr	Ile 1289		Thr	Gly	Ala	Pro 1290		Thr	Туг	Ser	Thr 1295	
30	G-1 y	Lys	Phe	Leu 1300		Asp	Gly	Gly	Cys 1305		Gly	Gly	Ala	Туг <b>13</b> 10		Ile
	Ile	Ile	Cys 1315		G1u	Cys	His	Ser 1320		Asp	Ser	Thr	Thr 1325		Leu	Gly
35	Ile	G1y 1330		Val	Leu	Asp	Gln 133	Ala 5	Glu	Thr	Ala	Gly 1340		Arg	Leu	Val
	Val 134		Ala	Thr	Ala	Thr 1350		Pro	Gly	Ser	Val 1355		Val	Pro	His	Pro 1360
40	Asn	Ile	Glu	Glu	Va1 136		Leu	Ser	Asn	Thr 1370		G1u	Ile	Pro	Phe 137	
	Gly	Lys	Ala	Ile 1380		Ile	G1u	Ala	Ile 138		Gly	Gly	Arg	His 139		Ile
45	Phe	Cys	H1s 139		Lys	Lys	Lys	Cys 1400		Glu	Leu	Ala	A1a 140		Leu	Ser
	Gly	Leu 141		Ile	Asn	Ala	Val 141	Ala 5	Tyr	Tyr	Arg	Gly 1420		Asp	Val	Ser
50	Va1		Pro	Thr	Ile	Gly 143		Val	Val	۷a٦	Va 1		Thr	Asp	Ala	Leu 1440

10	Met	Thr	Gly	Tyr	Thr 1445		Asp	Phe	Asp	Ser 1450		Ile	Asp	Cys	Asn 1455	
	Cys	Va1	Thr	Gln 1460		Val	Asp	Phe	Ser 1465		Asp	Pro	Thr	Phe 1470		Ile
15	Glu	Thr	Thr 1475		Val	Pro	Gln	Asp 1480		Va1	Ser	Arg	Ser 1485		Arg	Arg
	Gly	Arg 1490		G∃y	Arg	Gly	Arg 1495	Arg	Gly	He	Tyr	Arg 1500		Va1	Thr	Pro
20	G1y 1508		Arg	Pro	Ser	Gly 1510		Phe	Asp	Ser	Ser 1515		Leu	Cys	Glu	Cys 1520
	Tyr	Asp	Ala	Gly	Cys 1525		Trp	Tyr	Glu	Leu 1530		Pro	A1a	Glu	Thr 1535	
25	۷a۱	Arg	Leu	Arg 1540		Tyr	Leu	Asn	Thr 1545		Gly	Leu	Pro	Va 1 1550		Gln
	Asp	His	Leu 155		Phe	Тгр	Glu	Ser 1560		Phe	Thr	Gly	Leu 156	Thr	His	Ile
30	Asp	Ala 1571		Phe	Leu	Ser	Gln 157!	Thr 5	Lys	Gln	Ala	Gly 1580		Asn	Phe	Pro
	Tyr 158		۷a۱	Ala	Туг	G1n 1590		Thr	Val	Cys	Ala 1595		Ala	Gln	Ala	Pro 1600
35	Pro	Pro	Ser	Trp	Asp 160		Met	Trp	Lys	Cys 161		Ile	Arg	Leu	Lys 161	
	Thr	Leu	His	Gly 1620		Thr	Pro	Leu	Leu 162		Arg	Leu	Gly	Ala 1636		G1n
40	Asn	G1u	Val 163		Leu	Thr	His	Pro 164		Thr	Lys	Tyr	11e 164	Met 5	Ala	Cys
	Met	Ser 165		Asp	Leu	Glu	Val 165	Va1 5	Thr	Ser	Thr	Trp 166	Val	Leu	Val	Gly
45	Gly 166		Leu	Ala	Ala	Leu 1670		Ala	Tyr	Cys	Leu 167		Thr	Gly	Ser	Val 1680
	Val	Ile	Val	Gly	Arg 168		Ile	Leu	Ser	Gly 169	Arg O	Pro	Ala	Ile	Val 169	Pro 5
50	Asp	Arg	G1u	Leu 170		Tyr	Gln	G1u	Phe 170	Asp 5	Glu	Met	G1u	G1u 171	Cys 0	Ala

10																
	Ser	His	Leu   1715	Pro	Tyr	Ile		G1n <sup>.</sup> 1720		Met	Gln	Leu	Ala 1725		Gln	Phe
15		G1n 1730	Lys /	Ala	Leu	Gly	Leu 1735		Gìn	Thr	Ala	Thr 1740		G1n	Ala	Glu
	Ala 1745		Ala i	Pro	Val	Val 1750		Ser	Lys	Trp	Arg 1755		Leu	Glu	Thr	Phe 1760
20	Trp	Ala	Lys !	His	Met 1765		Asn	Phe	Ile	Ser 1770		Ile	Gln	Tyr	Leu 1775	Ala
	Gly	Leu	Ser	Thr 1780		Pro	Gly	Asn	Pro 178		Ile	Ala	Ser	Leu 1790	Met )	Ala
	Phe	Thr	Ala 1795		Ile	Thr	Ser	Pro 1800		Thr	Thr	Gln	Ser 1805	Thr	Leu	Leu
25	Phe	Asn 1810	Ile D	Leu	Gly	Gly	Trp 1815		Ala	Ala	Gln	Leu 1820	Ala )	Pro	Pro	Ser
	A1a 1825		Ser	Ala	Phe	Val 1831		Ala	Gly	Ile	Ala 183	G1y 5	Ala	Ala	Val	Gly 1840
30	Ser	Ile	Gly	Leu	Gly 184		Va1	Leu	Va'l	Asp 185		Leu	Ala	Gly	Tyr 185	Gly 5
	Ala	Gly	Val	Ala 1860		Ala	Leu	Va1	Ala 186		Lys	۷a۱	Met	Ser 187	Gly O	Glu
35	Met	Pro	Ser 1875		Glu	Asp	Leu	Val 188		Leu	Leu	Pro	Ala 188	Ile 5	Leu	Ser
	Pro	Gly 189	Ala O	Leu	Val	Val	Gly 189		Val	Cys	Ala	A1a 190	Ile O	Leu	Arg	Arg
40	H1s 190		Gly	Pro	G1y	61u 191		Ala	۷a۱	Gln	191	Met 5	Asn	Arg	Leu	Ile 1920
	Ala	Phe	Ala	Ser	Arg 192		Asn	His	۷a۱	Ser 193	Pro	Thr	His	Tyr	Va) 193	Pro 5
45	Glu	Ser	Asp	Ala 194		Ala	Arg	Val	Thr 194		ılle	Leu	Ser	Ser 195	leu O	Thr
	Ile	Thr	G1n 195		Leu	Lys	Arg	Leu 196		Gir	1 Trp	Ile	196	G1 և 5	Asp	Cys
. 50	Ser	Thr 197	Pro	Cys	Ser	Gly	Ser 197	Trp 5	Let	Arg	g Asp	Val	Trp 0	Asp	Trp	Ile

	Cys 1985		Val	Leu	Thr	Asp 1990	Phe	Lys	Thr	Trp	Leu 1995	Gln -	Ser	Lys	Leu	Leu 2000
10	Pro (	G1.n	Leu	Pro	Gly 2005		Рго	Phe	Phe	Ser 2010	Cys	Gln	Arg	Gly	Tyr 2015	Lys
	Gly	Val	Тгр	Arg 2020		ÁSÞ	Gly	Ile	Met 2025	G1n	Thr	Thr	Cys	Рго 2030	Cys	G1y
15	Ala	Gln	Ile 2035		G1y	His	Va∕l	Lys 2040	Asn )	Gly	Ser	Met	Arg 2045	Ile	Val	Gly
20	Pro	Lys 2050		Cys	Ser	Asn	Thr 205		His	Gly	Thr	Phe 2060	Pro	Ile	Asn	Ala
	Tyr 2065		Thr	Gly	Pro	Cys 2070	Thr	Pro	Ser	Pro	A1a 2075	Pro	Asn	Туг	Ser	Arg 2080
25	A1·a	Leu	Trp	Arg	Va1 208		Ala	Glu	Glu	Tyr 209	<b>Val</b> 0	Glu	Val	Thr	Arg 209	Val
	Gly	Asp	Phe	His 210		Val	Thr	Gly	Met 210	Thr 5	Thr	Asp	Asn	Va1 211	Lys 0	Cys
30	Pro	Cys	Gln 211		Pro	Ala	Pro	G1u 212	Phe 0	Phe	Ser	Glu	Val 212	Asp 5	G1y	Val
	Arg	Leu 213		Arg	Туг	Ala	Рго 213	A1a 5	Cys	Arg	Pro	Leu 214	Leu O	Arg	Glu	Glu
35	Val 214		Phe	Gln	Val	G1y 215		Asn	Gln	Tyr	Leu 215	Va1 5	Gly	Ser	Gin	Leu 2160
	Pro	Cys	Glu	Pro	G1u 216		Asp	Val	Ala	Va1 217	Leu 0	Thr	Ser	Met	Leu 217	Thr 5
40	Asp	Pro	Ser	H15		Thr	Ala	G1t	1 Thr 218		lys	Arg	Arg	219	Ala O	Arg
	Gly	Ser	Pro 219		Ser	Leu	Ala	Ser 220	- Ser	Ser	· Ala	Ser	G]r 220	leu 15	ı Ser	Ala
45	Pro	Ser 221		ı Ly:	s Ala	1 Thr	Cys	s Thi	r Thr	- His	s His	222	Sei 20	Pro	ASF	Ala
	Asp 222		ı Ile	e Gli	sfA u	AST 223	Lei 30	ı Le	յ ⊺րդ	o Ar	g G1r 223	1 G11 35	ı Me1	: Gly	/ G1)	/ Asn 2240
50	Ile	• Th	r Arg	y Va		u Ser 45				s Va 22	7 Va <sup>.</sup> 50	l Va	Le	ı Ası	Ser 22!	Phe

10	Asp	Pro		Arg 2260		Glu	G <sub>.</sub> 1u	Asp	G1u 2265	Arg	Glu	Val	Ser	Va1 2270	Pro .	Ala
	Glu	Ile	Leu 2275		Lys	Ser	Lys	Lys 2280	Phe	Pro	Ala	afA	Met 2285	Pro	IJe	Trp
15	Ala	Arg 2290		Asp	Туг	Asn	Pro 229	Pro	Leu	Leu	G1u	Ser 2300	Trp	Lys	Asp	Pro
	Asp 2305	Tyr	Val	Pro	Pro	Va1 2310	Val )	His	Gly	Cys	Pro 2315	Leu	Pro	Pro	Ile	Lys 2 <b>320</b>
20	Ala	Pro	Pro	Ile	Pro 232	Pro S	Pro	Arg	Arg	Lys 2330	Arg D	Thr	Val	Val	Leu 2335	Thr
	G1u	Ser	Ser	Va 1 234		Ser	Ala	Leu	A1a 234	Glu 5	Leu	Ala	Thr	Lys 2350	Thr )	Phe
25	Gly	Ser	Ser 235		Ser	Ser	Ala	Va1 236	Asp O	Ser	Gly	Thr	A1a 236	Thr 5	Ala	Leu
	Pro	Asp 237		Ala	Ser	Asp	Asp 237	G1y 5	Asp	Lys	Gly	Ser 238	Asp 0	Val	Glu	Ser
30	Tyr 238		Ser	Met	. Рго	Pro 239		G Tu	Gly	Glu	Pro 239	G1y 5	Asp	Pro	Asp	Leu 2400
	Ser	Asp	Gly	' Ser	Trp 240	Ser	Thr	- Val	Ser	G1u 241	1 61u 10	ı Ala	Ser	Glu	Asp 241	Val 5
35	Va1	l Cys	Cys	Ser 242	r Met 20	Ser	Туі	r Thr	Trp 242	Thr 5	- Gly	Ala	Leu	1 le 243	Thr 0	Pro
	Çys	s A1:	a Ala 24		u G1u	ı Sei	Ly	s Let 244	ı Pro	ıle	e Asr	n Ala	244	ı Ser 15	· Asn	Ser
40	Le	u Lei 24		g Hi	s His	s Ası	Me <sup>*</sup>	t Va: 55	Туг	- Ala	a Thi	r Thi	- Sei 60	- Arg	Ser	Ala
	G1 24		u Ar	g G1	n Ly:	s Ly 24	s Va 70	1 Th	r Phe	e As	p Ar	g Lei 75	u Gli	n Va`	i Leu	1 ASP 2480
45	As	řH q	s Ty	r Ar	g As 24	p Va 85	1 Le	u Ly:	s G1:	и Ме 24	t Ly: 90	s Al	a Ly:	s Ala	249	- Thr 95
	Va	l Ly	s Al	a Ly 25	s Le	u Le	u Se	r Va	1 G1 25	u G1 05	u Al	a Cy	s Ly	s Le 25	u Th 10	г Рго
50	Pr	o Hi	s Se	ir A1 515	a Ly	s Se	r Ly	's Ph 25	e G1 20	у Ту	r G1	y A1	a Ly 25	s As 25	p Va	1 Arg

•																
		Leu 2530		Ser	Lys	Ala	Va1 2535	Asn	His	Ile	His	Ser 2540	Val	Trp	Lys	Asp
10	Leu 2545		Glu	Asp	Thr	Val 2550	Thr	Pro	Ile	Asp	Thr 2555	Thr	Ile	Met	Ala	Lys 2560
	Asn	Glu	۷al		Cys 2565		Gìn	Pro	Glu	Lys 2570	G1y	Gly	Arg	Lys	Pro 2575	Ala
15	Arg	Leu	Ile	Va1 2580		Pro	Asp	Leu	Gly 2585	Val	Arg	Val	Cys	G1u 2590	Lys	Met
	Ala	Leu	Tyr 2595		Val	Val	Ser	Thr 2600	Leu )	Pro	Gln	Val	Val 2605	Met	Gly	Ser
20	Ser	Tyr 2610		Phe	Gln	Tyr	Ser 2615	Pro	Gly	Gln	Arg	Val 2620	G1u )	Phe	Leu	Val
25	Asn 262		Ťrp	Lys	Ser	Lys 263	Lys 0	Asn	Pro	Met	Gly 263	Phe 5	Ser	Tyr	Asp	Thr 2640
	Arg	Cys	Phe	Asp	Ser 264		Val	Thr	Glu	Asn 265	Asp 0	Ile	Arg	Val	G1u 265	Glu 5
30	Ser	Ile	Tyr	G1n 266		Cys	Asp	Leu	A1a 266	Pro 5	Glu	Ala	Arg	G1n 267	Ala D	Ile
	Lys	Ser	Leu 267		Glu	Arg	Leu	Tyr 268	Ile O	Gly	G1y	Pro	Leu 268	Thr 5	Asn	Ser
35	Lys	G1y 269		Asn	Cys	Gly	7yr 269	Arg 5	Arg	Cys	Arg	A7a 270	Ser O	Gly	Val	Leu
	Thr 270		Ser	Cys	G1y	Asr 271		Leu	. Thr	· Cys	7yr 271	Leu 5	Lys	Ala	Ser	Ala 2720
40	Ala	ı Cys	s Arg	A-1a	A 1 8		s Leu	Glr	ı Asp	273	5 Thr 30	Met	. Leu	Va1	Asn 273	Gly 5
	Asp	) Ast	Leu	y Va1		I I le	a Cys	; G1:	5ei 27	- A18 45	a Gly	/ Thr	- Glr	G1u 275	Asp 0	Ala
45	Ala	a Sei	r Let 27		y Va	l Phe	e Thi	- G10 271	ı Ala	a Mei	t Th	r Arg	7 Tyr 276	- Ser 35	- A1a	Pro
50	Pro	0 G1:		Pro	) Pro	G G T	n Pro 27	G G 10 75	и Ту	r As	p Le	u G1u 278	ı Lei 30	ı Ile	• Thi	Ser
50		s Se 85	r Se	r As	n Va	1 Se 27	r Va 90	1 A1	a Hi	s As	p Al 27	a Sei 95	r Gly	y Lys	s Ar	y Val 2800

10	Туг	Tyr	Leu	Thr	Arg 2805		bio	Thr	Thr	Pro 2810	Leu	Ala	Arg	Ala	Ala 2815	Trp
10	GTu	Thr	Ala	Arg 2820	His	Thr	Pro	Val	Asn 2825	Ser	Trp	Leu	Gly	Asn 2830	Ile	lle
15	Met	Tyr	Ala 2835		Thr	Leu	Trp	Ala 2840	Arg	Met	Ile	Leu	Met 2845	Thr	His	Phe
	Phe	Ser 2850		Leu	Leu	Ala	G1n 2855		Gln	Leu	Glu	Lys 2860	Ala )	Leu	Asp	Cys
20	G1n 286		Tyr	Gly	Ala	Cys 2870	Tyr )	Şer	Ile	Glu	Pro 2875	Leu	Asp	Leu	Pro	G1n 2880
	Ile	Ile	Glu	Агg	Leu 288		Gly	Leu	Ser	A1a 2890	Phe	Ser	Leu	His	Ser 2895	Туг
25	Ser	Pro	Gly	G1u 290	Ile 0	Asn	Arg	۷a۱	A1a 290		Cys	Leu	Arg	Lys 2910	Leu )	G1y
	۷a۱	Pro	Pro 291		Arg	۷a۱	Trp	Arg 292	His O	Arg	Ala	Arg	Ser 292	Val 5	Arg	Ala
30	Arg	Leu 293		Ser	Gln	Gly	Gly 293	Arg 5	Ala	Ala	Thr	Cys 294	G1y 0	Lys	Tyr	Leu
	Phe 294		Trp	Ala	Val	Lys 295		Lys	Leu	Lys	Leu 295	Thr 5	Pro	Ile	Pro	Ala 2960
35	Ala	Ser	Arg	Leu	Asp 296		Ser	Gly	Тгр	Phe 297	Val D	Ala	Gly	Tyr	Ser 297	Gly 5
	Gly	/ Asp	ıle	7yr 298	His	Ser	Leu	Ser	Arg 298	A 1 a	Arg	Pro	Arg	7rp 299	Phe 0	Met
40	Leu	ı Cys	299		. Leu	Leu	Ser	Va 1	G1y 10	/ Val	Gly	Ile	300	Leu 15	Leu	Pro
	Asr	301	-													
45																

#### Claims

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- An isolated non-A, non-B hepatitis virus particle comprising at least one antigen selected from the group consisting of a core antigen, a matrix antigen and an envelope antigen of the non-A, non-B hepatitis virus.
  - 2. The non-A, non-B hepatitis virus particle according to claim 1, wherein said core antigen, matrix antigen and envelope antigen are, respectively, coded for by a nucleotide sequence of the 333rd to 677th nucleotides, a nucleotide sequence of the 678th to 905th nucleotides and a nucleotide sequence of the 906th to 1499th nucleotides of the non-A, non-B hepatitis virus entire nucleotide sequence from the 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2 (16) hereof.
  - The non-A, non-B hepatitis virus particle according to claim 1 or 2, which has a ribonucleic acid corresponding to at least part of the nucleotide sequence shown in Fig. 2(1) through Fig. 2(16) hereof.
    - 4. A method for producing an isolated non-A, non-B hepatitis virus particle, which comprises:
      - (a) providing not more than ten different cDNA clones each comprising at least 1000 nucleotides and prepared from a non-A, non-B hepatitis virus genomic RNA fragment of at least 1000 nucleotides, said not more than ten different cDNA clones containing their respective cloned cDNA fragments which, on the whole, cover a region of at least the 333rd to 5177th nucleotides of the non-A, non-B hepatitis virus entire nucleotide sequence from the 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2(16) hereof:
- (b) taking out said cDNA fragments from said cDNA clones by cutting so as to respectively have predetermined nucleotide sequences such that when the predetermined nucleotide sequences are arranged in sequence, the resultant nucleotide sequence has at least a region which coincides with the region of the 333rd to 5177th nucleotides;
  - (c) ligating said taken-out cDNA fragments respectively having said predetermined nucleotide sequences in sequence to thereby construct a first deoxyribonucleic acid comprising a nucleotide sequence comprising at least the 333rd to 5177th nucleotides of the non-A, non-B hepatitis virus entire nucleotide sequence from the 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2(16) hereof;
  - (d) introducing at least one deoxyribonucleic acid selected from said first deoxyribonucleic acid and a second deoxyribonucleic acid obtained by substituting at least one nucleotide of the nucleotide sequence of said first deoxyribonucleic acid in accordance with the degeneracy of the genetic code into a replicable expression vector selected from a plasmid and an animal virus gene to obtain a replicable recombinant DNA comprising said plasmid and said at least one deoxyribonucleic acid introduced therein when said replicable expression vector is a plasmid or obtain a recombinant virus comprising said animal virus and said at least one deoxyribonucleic acid introduced therein when said replicable expression vector is an animal virus gene;
  - (e) transfecting prokaryotic or eukaryotic cells with said recombinant DNA when said replicable expression vector used in step (d) is a plasmid, to thereby form a transformant, followed by selection of said transformant from parent cells of the prokaryotic or eukaryotic cell culture;
    - (f) culturing said transformant obtained in step (e) in prokaryotic or eukaryotic cells to thereby produce a non-A, non-B hepatitis virus particle, or culturing said recombinant virus obtained in step (d,) in eukaryotic cells to thereby produce a non-A, non-B hepatitis virus particle together with an animal virus;
    - (g) isolating said non-A, non-B hepatitis virus particle.
  - 5. The method according to claim 4, wherein said first deoxyribonucieic acid comprises a nucleotide sequence of the 333rd to 5918th nucleotides.
  - The method according to claim 4, wherein said first deoxyribonucleic acid comprises a nucleotide sequence of the 333rd to 6371st nucleotides.
- The method according to claim 4, wherein said first deoxyribonucleic acid comprises a nucleotide sequence of the 333rd to 9362nd nucleotides.
  - 8. The method according to claim 4, wherein said first deoxyribonucleic acid comprises a nucleotide sequ-

ence of the 1st to 9416th nucleotides.

- 9. A recombinant comprising a replicable expression vector selected from a plasmid and an animal virus gene and a deoxyribonucleic acid comprising at least one nucleotide sequence selected from the group consisting of a first nucleotide sequence of the 333rd to 5177th nucleotides of the non-A, non-B hepatitis virus entire nucleotide sequence from the 1st to 9416th nucleotides shown in Fig. 2(1) through Fig. 2(16) hereof and a second nucleotide sequence obtained by substituting at least one nucleotide of said first nucleotide sequence in accordance with the degeneracy of the genetic code.
- 10. The recombinant according to claim 9, wherein said first nucleotide sequence comprises a nucleotide sequence of the 333rd to 5918th nucleotides.
  - 11. The recombinant according to claim 9, wherein said first nucleotide sequence comprises a nucleotide sequence of the 333rd to 6371st nucleotides.
  - 12. The recombinant according to claim 9, wherein said first nucleotide sequence comprises a nucleotide sequence of the 333rd to 9362nd nucleotides.
- The recombinant according to claim 9, wherein said first nucleotide sequence comprises a nucleotide
   sequence of the 1st to 9416th nucleotides.
  - 14. A diagnostic agent for the detection of non-A, non-B hepatitis by an antigen-antibody reaction, comprising an effective amount, for the antigen-antibody reaction, of the non-A, non-B hepatitis virus particle according to claim 1 or 2.
- 15. A vaccine for non-A, non-B hepatitis, comprising an effective immunogenic amount of a non-A, non-B hepatitis virus particle according to claim 1 or 2, and at least one pharmaceutically acceptable carrier, diluent or excipient.
- 16. Escherichia coli strain BK102 carrying a non-A, non-B hepatitis virus genomic cDNA, deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3384.
  - Escherichia coli strain BK106 carrying a non-A, non-B hepatitis virus genomic cDNA, deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3385.
  - Escherichia coli strain BK112 carrying a non-A, non-B hepatitis virus genomic cDNA, deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3386.
  - Escherichia coli strain BK146 carrying a non-A, non-B hepatitis virus genomic cDNA, deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3387.
    - Escherichia coli strain BK147 carrying a non-A, non-B hepatitis virus genomic cDNA, deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3388.
- Escherichia coli strain BK157 carrying a non-A, non-B hepatitis virus genomic cDNA, deposited at Fermentation Research Institute, Japan under the accession number FERM BP-3243.

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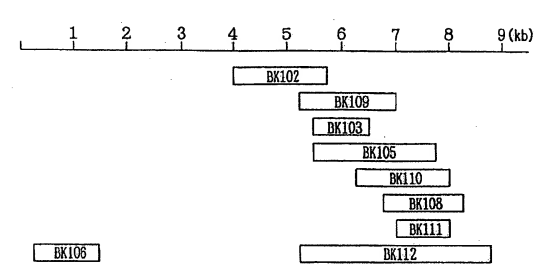
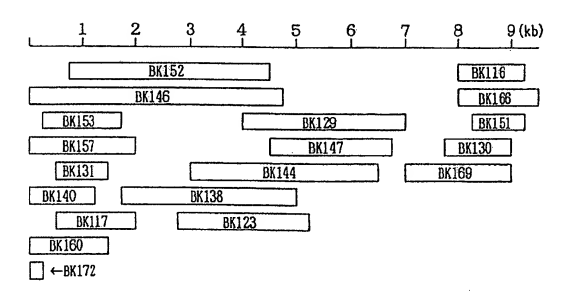


FIG. 1(2)



# FIG. 2(1)

CEATTGGCGACACTCCACCATAGATCACTCCCCTGTGAGGAACTACTGTCTTCAGGCGAAAGCGTCTAGCCATGGCGTTAGTATGAGTGTGGTGTGCAGCCTCCAGGACCCCCTCC GCTAACCCCCCCTGTGAGGTGGTATCTAGTGAGGGGACACTCCTTGATGACAGAGGTGCGTCTTTGGCAGTACGGCAATCATACTCACAGCAGGTCGTGGAGGTCCTGGGGGGAGG GGGAGAGCCATAGTGGT CTGCGGAACCGGTACACCGGAATTGCCAGGACGACGACGTCCTTT CTTGGATCAACCGGCTCAATGCCTGGAGATTTGGGCTGCCCCGGGAGACTG 121

MetSerThrAsnProLysProGlnArgLys CTAGECCIAGTAGTGTTGGGAAAAGGCCTTGTGGTACTGCCTGATAGGGTGCTTGCGAGTGCCCCCGGGGGGGACGTGCGTAGACCTTGAAAGGTCTAAAGGTCAAAAGAA GATCHSCT CATCACAACCCAGGGTTTCCGGAACACCATGACGGACTATCCCACGAACGCTCACGGGCCCTCCAGAGCATCTGGGACGTGGTACTCGTGCTTAGGATTTGGAGTTTCTT 241

ThrLysArgAsnThrAsnArgArgProGlnAspValLysPheProGlyGlyGlyGlnIleValGlyGlyValTyrLeuLeuProArgArgGlyProArgLeuGlyValArgAlaProArg 361

LysThrSerGludrgSerGlnProdrgGlyArgArgGlnProIJeProLysAlaArgArgProGluGlyArgThrTrpAlaGlnProGlyTyrProTrpProLeufyrGlyAsnGluGly CCTT OTGAAGG CT CGCCAGOCTT GOOD OF THE CONTRACT AGGG ST TO CAGGG COCCATION OF THE CONTRACT OF THE CONTRACT ACCOUNT GGAAGACTT CCGAGCGGT CGCCACCT CGTGGAAGGCGAAACCT AT CCCCAAGGCT CGCCGGGCCCGAGGCCAGGACT CGGCCCGGGGTACCCTTTGGCCT CTCTATGGCAATGAGG 481

## FIG. 2(2)

LeuGlyTrpAlaGlyTrpLeuLeuSerProArgGlySerArgProSerTrpGlyProThrAspProArgArgArgSerArgAsnLeuGlyLysYal1leAspThrLeuThrCysGlyPhe GCTT AGGSTGGGCAGGATGGCTCCTGTCACCCCGCGGGCTCCCGGCCCCACGGGCCCCCACGGACCCCCGGGGTAGTTGGGTAATTTGGGTAAGGTCATCGATACCCTCACATGCGGCT GAATCCACCOTCCTACCAAGACAGTGGGGCGCCGGATCAACCCCGGGTGCCTGGGGCCGCATCCAGGCATTAAACCCATTCCAGTAGGTATGGAGTGTAAGGCCGA 601

AlaAspLeuMetGlyTyrIleProLeuValGlyAlaProLeuGlyGlyAlaAlaArgAlaLeuAlaHisGlyValArgValLeuGluAspGlyValAsnTyrAlaThrGlyAsnLeuPro AGOGGCT AGAGT ACCUCATGT AAGGCGAGCAGCCGCGGGGACCCCCCGGGGACGGTCCCGGGACCGTGTA CCAGGCCCCAAGACCTCCTGCCGCGCACTTGATACGTTGTCCCTTAGACG 721

GlyCysSerPheSerIlePheLeuLeuAlaLeuLeuSerCysLeuThrThrProAlaSerAlaTyrGluValHisAsnValSerGlyIleTyrHisValThrAsnAspCysSerAsnAla 8

SerI leValTyrGluAlaAspLeuI leMetHisThrProGlyCysValProCysValArgGluGlyAsnSerSerArgCysTrpValAlaLeuThrProThrLeuAlaAlaArgAsn CAAGCATTGTGTATGAGGCAGCACTTGATCATGCATACTCCTGGGTGCCGTGCCTGCGTTCGGGAAGGCAACTCCTCCCTGGTGGGGTAGCGCTCACTCCCACGCTGGCGCAGCCAGG GTTOTTAACACATACTCOSTOGOCTGAACTAGTAOGTATGAGGACCCAOGCCAGGCCCTTOOGTTGAGGAGGGCGAAGGCCATOGOGAGTGAGGGTGGAAGGTGGAGGTCCT 961

ValThrIleProThrThrIleArgArgHisValAspLeuLeuValGlyAlaAlaAlaAlaPheCysSerAlaMetTyrValGlyAspLeuCysGlySerValPheLeuValSerGInLeu TGCAGTGGTAGGGGTGGTGCTGCTGCGGGTGCAGCTAGAGGAACCCCGGCCGACGAAAGACAAGGCATGCACCCCTGGAGAGACGACTAGACAAAAAGAAGAAGAGAGTCG 1081

## FIG. 2(3)

PheThrPheScrProArgArgIIIsVaIThrLeuGlnAspCysAsnCysScrIleTyrProGlyIIIsVaISerGlyIIIsArgMetAlaTrpAspMetMetMetAsnTrpSerProThrThr ACAMETIGGAAGAGGGGGGGGGGGTACACTGTATGTCCTGACATTGACGAGTTAAATAGGGCGGTACAGCCACTGGCATACGGAACGCTGTACTACTACTTGACCAGGGGGTGTT 1201

AlaLcuValValSerGInLcuLcuArgIleProGInAlaValValAspMctValAlaGlyAlaHisTrpGlyValLcuAlaGlyLcuAlaTyrTyrScrMctAlaGlyAsnTrpAlaLys CAGOOT ACTEGICIOCAGITACICOGGATCOCACAAGOOTOTIGGACATGGTGGOGGGGOOCACTGGGGAGTOOTGGOGGGCTTGOCTACTATTOCATGGOGGGAACTGGGCTA STOSSSATCACCACAGOSTCAATGAGGGGTGTTOGGCAGCACCTGTACCACGCCCCCGAGTGACCCCTCAGGACGGCCCGGAACAGATGATAAGGTACCGCCCTTGACCGAT 1321

ValLeuI leValMetLeuLeuPheAlaGlyValAspGlyAspThrHisValThrGlyGlyAlaGlnAlaLysThrThrAsnArgLeuValSerWetPheAlaSerGlyProSerGlnLys AGETTOTGATTGTGATGCTACTTTTTGCTGGGGTTTGACXGGGATACCCACGTGACXGGGGGGGGGCGAAAACCAACAAGGGGTGGTGTTCGCAAGTGGGGCGGTTGTTCG ICCAAGACTAACACTAGGATGAAAAAGGACGCCCAACTGCCCCTATGGGTGCTGTCCCCCCGGGTTGGGTTTGTGGTTGTCGGAGCACAGGTACAAGGGTTCACCGGGCAGAGTCT 1441

I leGInLeuI leAsnThrAsnGlySerTrpHisIleAsnArgThrAlaLeuAsnCysAsnAspSerLeuGInThrGlyPheLeuAlaAlaLeuPheTyrThrHisSerPheAsnSerSer ITTAGGTOGAATATITUSIGGITACCOTCAACOOTGTAGITIGICOTGAOGGGAOTTAACTGAGAGGGTOTGACOCAAAGAAOGGOGOGACAAGATGTGTGTATCAAGGTTGAGCA 1561

GlytysProGluArgMetAlaGlnCysArgThrIleAsplysPheAspGlnGlyTrpGlyProIleThrTyrAlaGluSerSerArgSerAspGlnArgProTyrCysTrpHisTyrPro COSSITECCLAGAGOSCATGSCCCAGTGCCACTTGACAAGTT CJACCAGGGATGGGGTTACTTATGCTGAGTCTAGCAGATCAGAGACCAGAGGCCATATTGCTGGCACATACC GECCCAOGGETCT CECCTACOGGETCACGCTGGTAACTGTTCAAGCTGGTCCCTACCCCAGGGTAATGAATACGACTCAGATGGTCTAGTCTGGTCTGGTATAAGGACGGTGATGG 1681

# FIG. 2(4)

ProProGinCysThrI leValProAlaSerGluValCysGlyProValTyrCysPheThrProSerProValValValGlyThrThrAspArgPheGlyValProThrTyrArgTrpGly CACCTCCACATGTACCATCTACCTCCTTCCTCCAGGTGTGCCCCCAGTGTACTCCTTCACCCCAGGCCCTGTCGTCGTCGGACCGATCGTTTCGGTGTCCCTACGTATAGATGGG STOSA OCTUBE TA CATOSTA GCATOSA OCA COCOCO COCOCO SOCIO CA CACA CASO OCA CASO A 1801

GludsnGluThrAspValLeuLeuLeuAsnAsnThrArgProProGlnGlyAsnTrpPheGlyCysThrTrpMetAsnSerThrGlyPheThrLysThrCysGlyGlyProProCysAsn GEGABAACEAGACTGACTGCTGCTCAACAACAACGCGCCCCCCCAAGAGCAACTGGTTCGCCTGCACATGGATGAATAGCACCGGGTTCACCAAGACATGTGGGGGGCCCCCCGTGTA CCCT CTT GCT CICACTGCA CGACGACTATTGT GCGCCGCCGCCGCGCGTT CCGTTGACCAAGCCGTCT ACT ACT ACT TAT CGTGGCTCCTGT ACACCCCCAAGCCGAC 1921

IlcGlyGlyValGlyAsnAsnThrLeuThrCysProThrAspCysPheArgLysHisProGluAlaThrTyrThrLysCysGlySerGlyProTrpLeuThrProArgCysHetValAsp ACAT CASCECISTICASCA ACACCOTG ACCOTG COTG COCCACACACOTG COTTCCCO ACCOCCACACACACA ACTOTG STOCATOS COCTAG COTTAG COT 2041

TyrProTyrArgLeuTrpHisTyrProCysThrYalAsnPheThrIlePheLysValArgMetTyrValGlyGlyValGluHisArgLeuAsnAlaAlaCysAsnTrpThrArgGlyGlu ACTATOCATACAGGOTOGGCATTACOCOTGCATOTTAACTTTACCATOTICAAGGTTAGGATGTATGIGGGGGGGGGGGGCACAGGCTCAATGCTGCATGCATGCATTGGACOCGAGGAG 2161

ArgCysAspLeuG1uAspArgProG1uLeuSerProLeuLeuLeuSerThrThrG1vTrpG1nVa1LeuProCysSerPheThrThrLeuProA1aLeuSerThrG1yLeu11e AGCCITICITES AGGACA GGGATA GGCCGGAGCT CAGCCCGCTGCTGCTGCTGCTACAACAA GAGTGGCAGGTACTGCCCTTCCTTCACCACCCTACCAGGCTCTGTCCACTGGCTTGA TOTO CACTE CACT CONTINUE CONTI 2281

## FIG. 2(5)

HisteulisGInAsn[leValAspValGInTyrLeuTyrGlyIleGlySerAlaValValSerPheAlaIleLysTrpGluTyrValLeuLeuLeuPheLeuLeuLcuCuAlaAspAlaArg TTCACCTCCATCAGAACATCGTGGACGTGCAATACGTATACGGTATAGGGTCAGCGGTTGTCTCTTTGCAATGAGAGTATGTGCGTGTTGCTTTTCCTTCTCGTAGGGAAGCAC  ValCysAlaCysLeuTrpMctMctLcuLcuLcuIleAlaGlnAlaGluAlaAlaLeuGluAsnLcuValValLcuAsnSerAlaSerValAlaGlyAlaHisGlyIleLcuSerPheLcuVal GTGT CTGTGCCTGCTTGTGGATGATGCTGCTGATAGCCCAGGCCCAAGCCCCTTGGAGAACCTGGTGGTCCTCAATTGGGCGTTGTGGCGGGGCGAAGGCATGCTTCCTTG 2521

PhePheCysAlaAlaTrpTyrIleLysGLyArgLenYalProGlyAlaThrTyrAlaLenTyrGLyYalTrpProLenLenLenLenLenAlaLenProProArgAlaTyrAlaMet TETTOTT CTGTGCCCCCCCCCCCCCAGAGGCAGGCTGGTCCCTGGGGCACATATGCTCTTTATGGCGTGTGCCGCCTGCTCCTGCTGCTGCCGCTTACCACCGGGAGCTTACCA NCANGAAGACACGGCGGACCATGTAGTTTCCCGTCCGACCCCCGCTGTATACGAGAATACCGCACACCGGCGACGAGGAGGAGAACACCATGAGGGCGCTCGAATGCGGT 2641

AspargGluMetAlaAlaSerCysGlyGlyAlaValPheValGlyLeuValLeuIeuThrLeuSerProTyrTyrLysValPheLeuAlaArgLeuIleTrpTrpLeuGlnTyrPheThr TGSACOGGGGGGGCGCGTGTGGGGGGGGGGGGGTTTTGGGGGTCTGGTACTCCTGACTTGTCACCATACTACAAGGTGTTCCTGGCTAGGCTCATATGGTTACAATATTTA ICOTGECOCTOTACCGAGGTAGCAGGCOTCCGGGCCAAAACACCCCAGACCATGAGGACTGAAAGTIGSTATGATGTTCCACAAGGAGGGGTCCCAGTATACCACCAATGTTATAAAAT 2761

ThrArgAlaGluAlaAspLeuHisValTrpIleProProLeuAsnAlaArgGlyGlyArgAspAlaIleIleLeuLeuMetCysAlaValHisProGluLeuIlePheAspIleThrLys 2881

#### FIG. 2(6)

Lculeul lehlal lelcuGlyProlcuMetValLcuGlnAlaGlyI leThrArgValProTyrMeValArgAlaGlnGlyLcuI leHisAlaCysMetLeuValArgLysValAlaGly 3001

GlyHisTyrValGlnMetAlaPheMetLysLeuGlyAlaLeuThrGlyThrTyrIleTyrAsnHisLeuThrProLeuArgAspTrpProArgAlaGlyLeuArgAspLeuAlaValAla 3121

ValCluProValValPheSerAspMetCluThrLysIleIleThrTrpGlyAlaAspThrAlaAlaCysGlyAspIleIleLeuGlyLeuProValSerAlaArgArgGlyLysGluIle 3241

LeuleuGlyProdlaAspSerleuGluGlyArgGlyLeuArgLeuLeuAlaProIleThrAlaTyrSerGlnGlnThrArgGlyLeuLeuGlyCysIleIleThrSerLeuThrGlyArg ITGAGGACCOGGCCCGGTATCAGAACTTCCCGCCCCCAACGCTGAGGAGGCGGAGTAGTGCCGGATGAGGTTGTCTGCGCCCCCGGATGAACCAACGTAGTGATGCAATGTCCGG TACTOCTEGECOGECOCATACTOTICA AGGOGGGGGGGTGOCATOCOTOGOCOCATOA OCACOCAACA GOGGGGOTA OTTGGTTGCATOA TOATACA GOGTTA CAGOTTA CAGOO 3361

AspLysAsnGlnValGluGlyGluValGlnValValSerThrAlaThrGlnSerPheLeuAlaThrOxsValAsnGlyValCysTrpThrValTyrHisGlyAlaGlySerLysThrLeu 3481

#### FIG. 2(7)

AlaAlaProLysGlyProIleThrGlnMctTyrThrAsnValAspGlnAspLcuValGlyTrpProLysProProGlyAlaArgSerLeuThrProCysThrCysGlySerSerAspLeu TAGCOSCICAMAGOGGICCAATCACCCAGATGTACACTAATGTGGACCAGGACCTGTCGGCTGGCCCAAGCOCCGGGGGGGGTTCCTTGACACCATGCACCTGTGGCAGCTCAGACC AT COSCOCIONATION CONTINUE TO SEGIONA A TOTA CONTROPO CONTROPO CONTINUE CONTROPO CONTINUE CONTINUE A CONTINUE C 3601

[yrLeuVa]ThrArgHisAlaAspVa]]]eProValArgArgAlyAspSerArgG]ySerLeuLeuSerProArgProValSerTyrLeuLysG]ySerSerG]yGJyProLeuLeu TTTACTTGGTCACGAGGATGGTGAGTCATTCCGGTGGGCGGGGAGAGGAGAGTAGGGGGGCTGCTCTCCCCAGGCCTGTGCTGCTTGAAGGGCTCTTGGGGGGTCTCACACTGC AATGAACCAGTGCTCTGTAGGACTGCAGTAAGGCCAGGGGGCGCCCCCCCTGTCATCCCCTGGAAGGAGGTTCCGGAGGAGGAGGAGGAGGAGGAGGAGGAGGCCACCAGGTGAGG 3721

CyspropheglyHisAlaValGlyIlePheArgAlaAlaValCysThrArgGlyValAlaLysAlaValAspPheValProValGluSerMetGluThrThrMetArgSerProValPhe l CTGCCCTT CGGCCACGCTGTGGGCTACTTCCCGGGTATGCACCCCGGGGGGTTGCGAAGGCGGTGGACTTTGTGCCCGTAGAGTCCATGGAAACTACTATGCGGTCTCC 3841

ThrAspAsnSerSerProProAlaValProGlnSerPheGlnValAlaHisLeuHisAlaProThrGlySerGlyLysSerThrLysValProAlaAlaTyrAlaAlaGlnGlyTyrLys TCACESACAACTCATCCCCCCCGGCCGTACCGCCATTTCAAGTGGCCCACCTACACGCTCCCACTGGCAAGGGCAAGAGTACTAAAGTGCCGGCCATATGCAGCCCAAGGGTACA AGT GCCT GTT GAGT AGGGGGCGGCCATGGCGT CAGTAAAGTT CACCGGGTGGATGTGCGGAGGTGACCTTCT CATGATTT CACGGCCGACGTATACGTCGCATGT 3961

ValLeuValLeuAsnProSerValAlaAlaThrLeuGlyPheGlyAlaTyrMetSerLysAlaHisGlyIleAspProAsnIleArgThrGlyValArgThrIleThrThrGlyAlaPro AGGICCT CGT CCT CAAT CCGT TG CCGCT ACCTT AGGGTTT GGGG CGTAT AT GT CT AAGG CA CCGT ATTG ACC CCAACAT CAGAACT GGGGT AAGGACCATT ACCACAGGCCCC 4081

#### FIG. 2(8)

ValThrTyrSerThrTyrGlyLysPhcLcuAlaAspGlyGlyCysSerGlyGlyAlaTyrAsp[leIleIleIleCysAspGluCysHisSerThrAspSerThrThrIleLeuGlyIleGly CECACTETATEAGATGEATACCETTCAAGAACCGCTACCACAAGACCCCCGGCAATACTCTAGTATTATACACTACTCACCGTAAGTTGACTGAGCTGATGTTAGAACCCGTAGTAGC ThrValLeuAspGlnAlaGluThrAlaGlyAlaArgLeuValValLeuAlaThrAlaThrProProGlySerValThrValProHisProAsnIleGluGluValAlaLeuSerAsnThr GCACNSTCCTGGACCANGGGGAGGAGGGGTGGGGGGGGTTGTGGTGGCCACGGGTAGGCTGGGGATGGGTCACGGGCACACACCCAAACATGAGGAGGTGGCCCTGTCTAATA OSTETCAGGA ECTGETT OSCOTOS COSACOTOS COGA A CAGO CAGA A GOGA A GOGA COCTA A GOCA A GOCA CAGA CAGA TOTO TA A CA 4321

GlyGluI]eProPheTyrGlyLysAlaI]eProI]eGluAlaIJeArgGlyGlyArgHisLeuI]ePheCysHisSerLysLysLysCysAspGluLeuAlaAlaLysLeuSerGlyLeu CTGGAGAGATCCCCTTCTATGGCAAAGCCATCCCCATTGAAGCCATCAGGGGGGGAAGGCATCTCATTTCTGTCATTCCAAGAAGAAGTGCGACGAGGCTCGCCGCAAAGCTGTCAAGGCC GACCICICICAGGGGAAGATACOGTITOGGTAGGGGTAACTICGGTAGTCCCCTICGGTAGAGTAAAGACAGTAAGGTTCTICAGGCTGCTCCAGGGGGGTTTTGGACAGTCCGG 4441

GlylleasnalayalalatyrtyrargGlyLeuaspyalSerVallleProThrIleGlyaspyalValValAlaThraspalaLeumetThrGlyTyrThrGlyAspPheAspSer TOGGATCAAGCIGTGGGGTATTACGGGGGGTGGATGTGTGCGGTCATACCAACTATGGAGGAGGTGGTTGTGGTGGCAACAGAGGCTGTGATGAGGGGGTATAGGGGGAGTTTGACT AGCTTAGTTGCGACACCGCATAATGGCCCCCCGAGCTACACAGGCCAGTATGGTTGATAGCCTCTGCAGCACACACCGTTGTCTGCGAGACTACTGCCCGATATGCCCGCTGAAACTGA 4561

ValileAspCysAsnThrCysValThrGlnThrValAspPheSerLeuAspProThrPheThrIleGluThrThrThrValProGlnAspAlaValSerArgSerGlnArgArgGlyArg CAGTGATCGACTGTAACACATGTGTCACCCGGACAGTGCAGGTTGGATCCCACCTTCACCATTGAGAGGAGGACGTGACGAGGGGGGGTGTGGGGGTTGGAGGGGGGTA GTCACTAGCTGACATTGTGTACACAGTGGGTCAGCTGAAGTGGAACCTAGGGTGGAAGTGGTAACTCTGCTGCTGGCAGGGAGGTTCTGGGTCACAGGGGGGAGGGTGGCGCCCCAT 4681

#### FIG. 2(9)

ThrClyArgGlyArgArgGly[leTyrArgPhcValThrProGlyGluArgProSerGlyMetPheAspSerSerValLeuCysGluCysTyrAspAlaGlyCysAlaTrpTyrGluLeu GGACTGGCAGGGT AGGAGGCATCT ACAGGTTTGTGACTCCGGGAGAACGCCCCTCGGGCATGTT CCATGCTCGTGTGTGTGTGTGTGTATGACGCGGGGTTTGCTTGGTACGAG CCTGACOCTICCCCATCCTCCOTAGATGTCCAAACACTGAGGCCCTCTTGCCGGGAGCCCGTACAAGGTAAAGGAGCGGGAGACAGGACTGCGGATACTGCGGGACACGAAACATGCTGG ThrProAlaGluThrSerValArgLeuArgAlaTyrLeuAsnThrProGlyLeuProValCysGlnAspHisLeuGluPheTrpGluSerValPheThrGlyLeuThrHisIleAspAla TCACCOCHECCEAGACCTCLEGTTGCGGGCCTACCTGAACACACCAGGGTTGCCCGTTTGCCAGGACCACCTGGAGTTCTGGGAGAGTGTCTTCACAGGCCTCACCCATATAGATG IGTEGGECOSICTCTGGAGCCAATCCAAGSCCGEGATGGACTTGTGTGTCCCAAGGGCAAAGGTCCTGGTGGAAGTCTCAAAAGTGTCGGGAGTGGGTATATCTAC HisPheLeuSerGlnThrLysGlnAlaGlyAspAsnPheProTyrLeuValAlaTyrGlnAlaThrValCysAlaArgAlaGlnAlaProProProSerTrpAspGlnMetTrpLysCys ITETGAAGAACAGGETCTGGTTCGTCCTCTCTGTTGAAGGGGATGGACCATGGTTCGGTGCCACAGCGGTCGGAGTCGGGGGTGGAGGTAGTACCCTAGTTTACACCTTCA 5041

Leul leargleulysProThrleuHisGlyProThrProLeuLeuTyrargLeuGlyAlaValGlnAsnGluValThrLeuThrHisProI]eThrLysTyrIleMetAlaCysMetSer  AlaAspLeuGluValValThrSerThrTrpValLeuValGlyGlyValLeuAlaAlaLeuAlaAlaTyrCysLeuThrThrGlySerValValIleValGlyArgIleIleLeuSerGly OSCITS ACTION ACTORICA CONTRACTION AND ACTION ASSESSED ASSESSED ASSESSED ASSESSED ASSESTED AS 3CCSACTIGSACTICCAGCAGTGATGTGGGCCCACCACCACCACCTCAGGAACCTGGGACCGGCATAAGGACTGTTGTCGGTCACCAGGACCATGGTAACACCCATGGTAA 5281

### FIG. 2(10)

ArgProAlaIleValProAspArgGluLcuLeuTyrGlnGluPheAspGluMetGluGluCysAlaSerHisLcuProTyrIlcGluGlnGlyMetGlnLeuAlaGluGlnPheLysGln GGAGGCCCGCCATTGTTCCCGACAGGGAGCTTCTCTACCAGGAGTTCGATGAAATGGAAGAGTGCCCCTCGCACCCCTTACATCGAGGAGTGCAGGGAATTCCAAGC CCTCDGCCGGTAACAAGGGCTGTCCCTCGAAGAATGGTCCTCAAGCTACTTTACCTTCTCAGGGAGCGTGGAGGGATGTAGCTCGTTCCCTTACGTCGAGGGGCTGTTAAGTTGG 5401

LysAlaLeuGlyLeuLeuGlnThrAlaThrLysGlnAlaGluAlaAlaAlaProValValGluSerLysTrpArgAlaLeuGluThrPheTrpAlaLysHisMetTrpAsnPheIleSer AGAAAGCICT COGGTT ACTGCAAACAGCCACCAAACAAGGGGGGGGGTGCTGCTGGTGGGGGTGGGGGGCCTTGAGACATTCTGGGGGAGCACGTGTGGAATTTCATCA 5521

GlyileGinTyrieuAlaGlyLeuScrThrLeuProGlyAsnProAlaIleAlaSerLeuMetAlaPheThrAlaSerIleThrSerProLeuThrThrGinSerThrLeuLeuPheAsn **GOGGGATACAGTACTTAGCAGGCTTATCCACTCTGCCTGGGAACCCGGCAATAGCATCATTGATGGCATTCACAGCCTCTCACAGCCCGCTCACCACAAGTACCCTCTGTTTA** 5641

I leLeuGlyGlyTrpValAlaAlaGlnLeuAlaProProSerAlaAlaSerAlaPheValGlyAlaGlyIleAlaGlyAlaAlaValGlySerIleGlyLeuGlyLysValLeuValAsp 5761

IleLeuAlaGlyTyrGlyAlaGlyValAlaGlyAlaLeuValAlaPheLysValMetSerGlyGluMetProSerThrGluAspLeuValAsnLeuLeuProAlaIleLeuSerProGly 5881

#### FIG. 2(11)

AlaLeuvalvalglyvalvalCysAlaAlaIleLeuvrgArgHisValGlyProGlyGluGlyAlaValGlnTrpMetAsnArgLcuIlcAlaPheAlaScrArgGlyAsnHisValScr GOCCCIGITOTOSGGICOTGIGIGGACATACIGOSIOSACACITISGICOSGAGAGAGAGACACIGICASTGAACOGGCIGATAGOSITOGCCIOGOGOGIAAICATITI **600** 

ProThrHisTyrValProGluSerAspAlaAlaAlaAlaArgValThrGlnIleLcuSerSerLcuThr1leThrGlnLcuLcuLysArgLcuHisGlnTrpIleAsnGluAspCysSerThr COCCCACICATITICOTICAGAGICACACICICAGAGICITACTCAGATICITACICAGCITTACATACATCACICACAGAGAGAGACACCACICAGAGATTAATISAAGACITICA IGGGTIGGT ICACKEACT CTCICICT GCGCGCGTCCCACAATGAGTCTAGGAGAGGTCGGAATGGTAGTGAGTCGACGACGTTTTCCCAGGTGGTCACCTAATTACTTCTGACGAGGT 6121

ProCysSerGlySerTrpLeuArgAspValTrpAspTrp1leCysThrValLeuThrAspPheLysThrTrpLeuGinSerLysLeuLeuProGlnLeuProGlyValProPhePheSer ITGGCACAGGCGGAGCGATTCCCTACAAACCCTGACCTATACSTGCCACAACTGAAGTTCTGGACGACGACGTCAGGTTCGAGGACGGCGTCCATGGACCTCAGGGAAAAAAGA CACCTIGITICCSSCTOTIGSCTAAGGGATGITITGGGACTIGSATATGCAQGGTTGACTGACTTCAAGACCTGGCTCCAGTCCAAGCTGCTGCCGGCGCAGCTACCTGGAGTCCCTTTTTTCT 6241

 $CysGInArgGIyTyrIysGIyValTrpArgGIyAspGIyI1eMetGInThrThrCysProCysGIyAlaGInI1eThrGIyHisValLysAsnGIySerMet\DeltargI1eValGIyProLysSInArgGIyTyrIysGIyValTrpArgGIyAspGIyI1eValGIyProLysSInArgGIyFinGIyHisValLysAsnGIySerMet\DeltargI1eValGIyProLysSInArgGIyFinGIyHisValLysAsnGIySerMet\DeltargI1eValGIyProLysSInArgGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyHisValLysAsnGIyFinGIyHisValLysAsnGIyHisValL$ OTICICAADICCIGITACAAGGGAAGTITIGGGGGAGAAGGAAGCACCTGCCCATGTGGAGGACACAGATCACDGACATGTCAAAAAGGTTCCATGAGGATGTTOGGGCGTA GCACSETTGCS CCCATGTT CCCTCAGACGCCCCTCTGCCGT AGT ACGTTTGGTGGACGGGT ACACT COTCTAGTCGCCTGT ACACTTTTT GCCAAGGT ACTCCTAGCAGCCCGA 6361

ThrCysSerAsnThrTrpHisGlyThrPheProIleAsnAlaTyrThrThrGlyProCysThrProSerProAlaProAsnTyrSerArgAlaLeuTrpArgValAlaAlaGluGluTyr ICTGGAOGTOSTTGTGCACOGTACCTTGTAAGGGGTATGOGTATGTGGOOGGGAOGTGTGGAGAGGTCGOGGTTTGATAAGATCCOGGGACACOGGCACOGGGACTCCTCA AGACCTGCAGCACACTGGCATGGAACATTCCCCATCAAGGCATACACCAGGGCCCCTGCACACCTCTCCAGGGCCAAACTAITCTAGGGGCTGTGTGGGGTGGCGGCTGAGGAGT 6481

ValGluValThrArgValGlyAspPhellisTyrValThrGlyMetThrThrAspAsnValLysCysProCysGlnValProAlaProGluPhePheSerGluValAspGlyValArgLeu TECACCTCCACTGCGCCCCAAAGGTGATGCACTGCCCGTACTGGTGACTGTTGCATTTCACGGGTAGGTCCAAGGCCGAGGACGTTAAGAAGAGCCTCCACCTGCCTCAGGCCA ACCIGEACEICACECIGEGEGEATITICEACTACEIGACGAIGACCACIGACAACGIAAAGIGCCCAIGCCAGGITCCASCICCIGAATICITICICAGGIGGACGAGGIGCAGI 6601

III SATGTYTA I APTOA I ACYSAT BPTOL EUL EUAT GGOUG IUYA I THTPHEGINYA I GIYL EUAS NGINTYTL EUVA I GIYSETGIN LEUPTOCYSGIU PTOA SIYA I AIA I AIA AIA TGCACAGGTAGGCTCCAGGCGTCCCTCCTAGGGAGGGTTACATTCCAGGTGGGGTCAACCAATACCTGGTTGGGTGACAGGTACCATGCGAGGCGGAACCGGATGTAGCAG NOTITICCATEGGAGECGGCAGGTCCGCAAGGATGCCCTCCAATGTAAGGTCCAAGCTGGTT ATGGACCAACCAGTGTGTGAGGTAGGTAGGTTGGCGTGGGCTTGGCCTACATGTTC 6721

LeuThrSerHetLeuThrAspProSerHisIleThrAlaGluThrAlaLysArgArgLeuAlaArgGlySerProProSerLeuAlaSerSerSerAlaSerGlnLeuSerAlaProSer TECTCACTTCCATGCTCACCCACCCTCCCACATCACAGAAAAGASCTAAGGSTAGGCTGCCAGGGGGTCTCCCCCTCGTTGGCCAGCTCTTCAGGTAGCCAGTTGTCTGGG 6841

LeulysAlaThrCysThrThrHisHisValSerProAspAlaAspLeuIleGluAlaAsnLeuLeuTrpArgGlnGluMetGlyGlyAsnIleThrArgValGluSerGluAsnLysVal GGAACTTCCGCTGTACGGGGGGAGGAGGCCTGCGAGTAGGTCCGGTTGGAGGACACGGCGTTCTTACCGGCCTTGTAGTGGGGGGCACCTCAGCCTCTTGTTCC CCTTGAAGGCGACATGCCACCATGTCTCTCCGGACGCTGACCTCATCGAGGCCAACCTCCTGTGGCGGCAGGAGATGGGCGGAGATCGGCGCGTGCACTGGAGTCGGAGTCGGAGTCGGAGTCGAACAAGG 6961

ValValLeuAspSerPheAspProLeuArgAlaGluGluAspGluArgGluYalSerValProAlaGluIleLeuArgLysSerLysLysPheProAlaAlaMetProIleTrpAlaArg ACCATCAGGACTCAGAAAAAGCTGGGGAAGCTGCCTGCTACTGTGCCTTTGATAGGCAAGGCGGCTGTAGGAGGCGTTTAGGTTGTTCAAGGGGGGGTGGTAGAGGGGG 7881

## FIG. 2(13)

ProdspTyrdsnProProLeuLeuGluSerTrpLysAspProdspTyrValProProValValHisGlyCysProLeuProProIleLysAlaProProIleProProArgArgLys GOCCIGATI ACAACCOT CCACTGT AGAGT CCTGGAAGGACCCGGACT ACGT CCCT CCGGTGGTGCACGT TGCCACCT AT CAAGGCCCCT CCAAT ACCACCT CCACGGAAAA OSGOCTAATUTTGGGAGGTGACAATCTCAGGACCTTCCTGGGGCTGATGCAGGCCACCACGTGCCCACGGGGCAACGGTGGATAGTTCCGGGGAGGTTATGGTGGAGGTGCCTCTT 7201

ArgThrValValLeuThrCluSerSerValSerSerAlaLeuAlaGluLeuAlaThrLysThrPheGlySerSerGluSerSerAlaValAspSerGlyThrAlaThrAlaLeuProAsp AGAGGACGGTTGTCCTAACAGAGTCCTCCGTGTTTCTGCGTTAGGGGAGGTGGGTACTAGGAGCTTCGGCAGCTCCGAATCATCGGCCGTGGACGGGGCACGGAGCGACCGTTCCTG 7321

GInAlaSerAspAspGlyAspLysGlySerAspYalGluSerTyrSerSerMetProProLeuGluGlyGluProGlyAspProAspLeuSerAspGlySerTrpSerThrYalSerGlu ACCAGGOCT COGACGATGACAAAGGAT COGACGITGAGTCGT ACT CCTCCATGCCCCCTTGAGGGGGAACCGGGGGACCCCGATCTCAGTGACGGGTCTTGGTCTACCGTGAGG TEST COSSAGE CIGCIGCO COTA GIOTICO TAGGOTO CAGO A TOAGGO GA A COSTA COSTA GOO COTAGA GOO CAGA GOO CAGA A COAGA TOGO COAGA GOO CAGA TOGO COAGA GOO CAGA TOGO COAGA GOO CAGA TOGO COAGA GOO CAGA A COAGA TOGO COAGA GOO CAGA A COAGA GOO CAGA A COAGA GOO CAGA A COAGA GOO CAGA A COAGA COAGA A 7441

GludlaSerGludspValValCysCysSerMetSerTyrThrTrpThrGlydlaLeuIleThrProCysAladlaGluGluSerLysLeuProIleAsnAlaLeuSerAsnSerLeuLeu 7561

ArghishisasnketvalTyralaThrThrSerargSeralaGlyLeuargGInLysLysValThrPheaspargLeuGInValLeuaspasphisTyrargaspYalLeuLysGluMet TGOSCCACCATAACATGSTTTATGCCACAACATCTCGCAGGCCGGCGGCAGAAGAAGGTCACCTTTGACAGACTGCAAGTCCTGGACGACCACTACCGGGACGTGCTCAAGGAGA ACICIETICETATICTACCAATACLEITICTAGAGCETCICICGGACGCCETCTTCTTCCAGTGGAAACTGTCTGACFTTCAGGACCTGCTGGTGATGGCCCTGCACGAGTTCCTCT 7681

### FIG. 2(14)

LysalaLysalaSerThrValLysLcuLeuSerValGluGluAlaCysLysLeuThrProProHisScrAlaLysScrLysPhcGlyTyrGlyAlaLysAspValArgAsnLeu TGAAGGOGAAGGOGTOCACAGTTAAAGGCTAAAACTCCTATCOGTAGAGGAAGCCTGCAAGCTGAOGCCOCACATTOGGCCAAATCCAAGTTTGGCTATGGGGCAAAGGAOGTCOGGAACC NETICOCTIO COCAGGI GI COM TITO COM AGGOAT AGGOAT CI COTIO COM COMO COMO TITA AGO TITA AGO TITO CA AGO A MANA M 7801

SerSeriysAlaValAsnHisIleHisSerValTrpLysAspLeuLcuGluAspThrValThrProIleAspThrThrIleMetAlaLysAsnGluValPheCysValGlnProGluLys TATECAGCAAGGEESTT AACEACATECAGTECGGTGTGGGAAGGACTTGGTAGACACTGTGACACCAATTGACACCACCATCATGGGAAAAAGAGAGTTTTGTGTGTACAACCAGAGA N AGET COTT COSECATTESTE AGET GAGG CACCTT CCT GAACGACCTT CTGT GACT GTG GGT GGT GGT GGT GGT ACT CTA AGA GAAGACACAGGT TGST CT CT 7921

GlyGlyArgLysProAlaArgLeuIleValPheProAspLeuGlyValArgValCysGluLysMetAlaLeuTyrAspValValSerThrLeuProGlnValValMetGlySerSerTyr AAGGAGGCCGTAAGCCAGCCCGTTATGTATTCCCCAGATCTGGGAGTCGTGTATGCGAGAAGATGGCCTCTATGATGTGGTGTGCCCTCCTCCGGGTGGTGTGATGGGGCTCCTCAT 8841

GlyPheGlnTyrSerProGlyGlnArgValGluPheLeuValAsnThrTrpLysSerLysLysAsnProMetGlyPheSerTyrAspThrArgCysPheAspSerThrValThrGluAsn IGCCTAAGGTCATGAGAGGACCOGTOGCTCAAGGACCACTTATGGACCTTTAGTTTTCTTTTTGGGGTACCOGAAAGTATACTGTGAGGACAAAGCTGAGTTGCCAGTGGCTCTCT 8161

Asp[leArgValGluGluSer]leTyrGlnCysCysAspLeuAlaProGluAlaArgGlnAlaIleLysSerLeuThrGluArgLeuTyrIleGlyGlyProLeuThrAsnSerLysGly AGGACAT COSTOTTGAGGAGT CAATTTACCAATGTTGACCTTGGCCCCGAAGCCAGACGGCCATAAAATGGCTCACAGAGGGGGGTTTATATGGGGGGTCCTGGACTAATTCAAAAG 8281

#### FIG. 2(15)

COSTOSTIGACICAATAGOGGCCACGGCGCCCACGCCCACGCCCACGCCATGCTGCAAGGCCATTGTGGCAAGGAACTTCCGGAGACGTCCGAACGCTCGAAGGTTTCGAGGTCCTGA GECALAACTECOGTTATCECCGGTGCCGCGCGCGCGTGCTGACGACTAGCTGCGGTAACCCCTCACATGTTGAAGECCTCTGCGAGCCTGTGAAGCTGCGAAGGTCCAGGACT <u>840</u>

 $Thr \texttt{MetLeu} \\ \textbf{MetMetMet} \\ \textbf{MethetMet} \\ \textbf{MethetMeth$ GCACCATATOCTICATAAAGAAGACCTCCTTATCTGTGAAAGCGCGGAACCCAAGAGGACGCGAGCGTACGAGGTGCTTCAGGGAGGTTATGACTAGGTACTCGGCCCCCC OT GCTA CLAGACITI GCCT CT GCT GGAGACATAGACA TAGACACT TT CGCGCCTT GGGTT CT CCTGCGCGCGCT CAGAAGTGCCT CCGATACTGATCCATGAGGCGGGGGGGG 8521

AspProProGInProGIuTyrAspLeuGluLeuIleThrSerCysSerSerAsnValSerValAlaHisAspAlaSerGlyLysArgYalTyrTyrLeuThrArgAspProThrThrPro GEEAUCOCICCCCAACCAGAATACEACTTGGAGCTGATAACATCATGTTCCTCCAATGTGTGGCCCAGGATGCATCAGGGAAAAGGGTGTACTACCTCACCGGTGATGACCACCACCA 8641

LeuAlaArgAlaAlaTrpGluThrAlaArgHisThrProValAsnSerTrpLeuGlyAsnIleIleMetTyrAlaProThrLeuTrpAlaArgMetIleLeuHetThrHisPhePheSer 8761

I leLeuLeuAlaGInGIuGInLeuGIuLysAlaLeuAspCysGInIIeTyrGIyAlaCysTyrSerIIeGIuProLeuAspLeuProGInIIeIIeGIuArgLeuHisGIyLeuSerAla 88

# FIG. 2(16)

PheSerLoullisSerTyrSerProGlyGluIleAsnArgValAlaSerCysLeuArgLysLeuGlyValProProLeuArgValTrpArgHisArgAlaArgLeu	)] CATITTCACTCCATAGITACTCCCACGTGAGATCAATAGGGTGGCTTCATGCGTCAGGAAACTTGGGGTACCACCTTGGGAGGTCTGGAAGAGATGAGAAAAAAAA	GTANA AGIGAGGTATCA A TCA CA CGTCCA CTCTA GTTA TCCA AGTA CGCA AGT TCTT TGAA CCCA TGGT CGG CA CGCT CA CA CCT CA CA CGCT CA CA CCT CA CA CGCT CA CA CGCT CA CA CGCT CA CA CGCT CA
	9001	

LeuSerGinGlyGlyArgAlaAlaThrCysGlyLysTyrLeuPheAsnTrpAlaYalLysThrLysLeuIhrProIJeProAlaAlaSerArgLeuAspLeuSerGlyTrPhe	Val AlaGLy TyrSer-GLyGLyAspAle TyrHisSerfeuSerArgAlaArgProArgTrpPheMetLeuCysLeuLeuLeuSerValGLyValGLyIleTyrLeuLeuProAsnArg
TACIGICCCAGGGAGGGAGGCCCCACTTGTGCCAAATACCTCTTCAACTGGACAAACCAAACTTAAACTCACTC	TOGTIOCTIOTTACACCOCCAGATATATCACACCTGTTCGTCCOCGACCCGTTGGTTCATGTCTTGCTACTTTCTGTAGGGTAGGCATCTACTGACATCTACTACTACTACTACTACTACTACTACTACTACTA

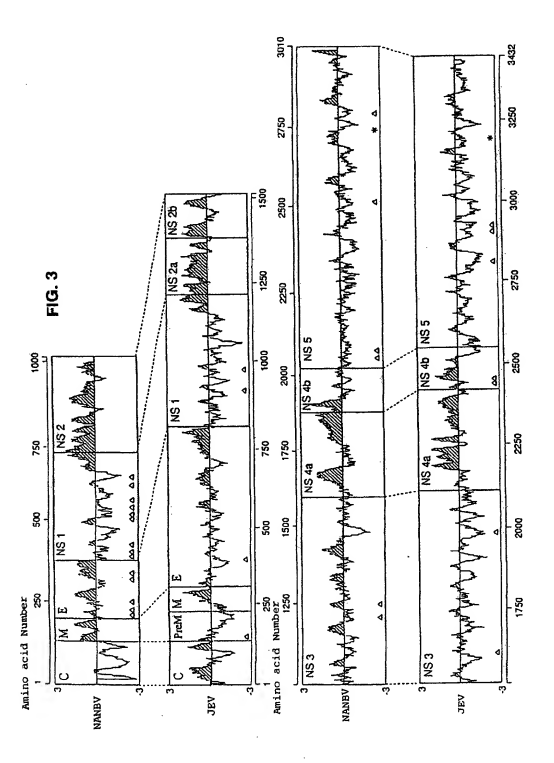


FIG. 4

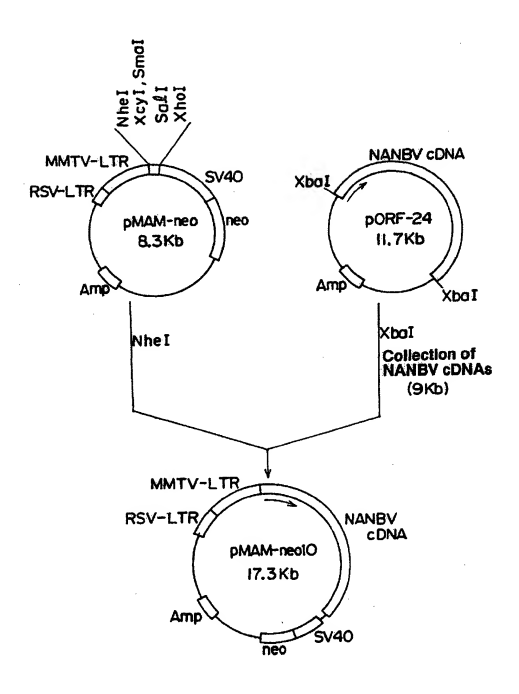


FIG. 5

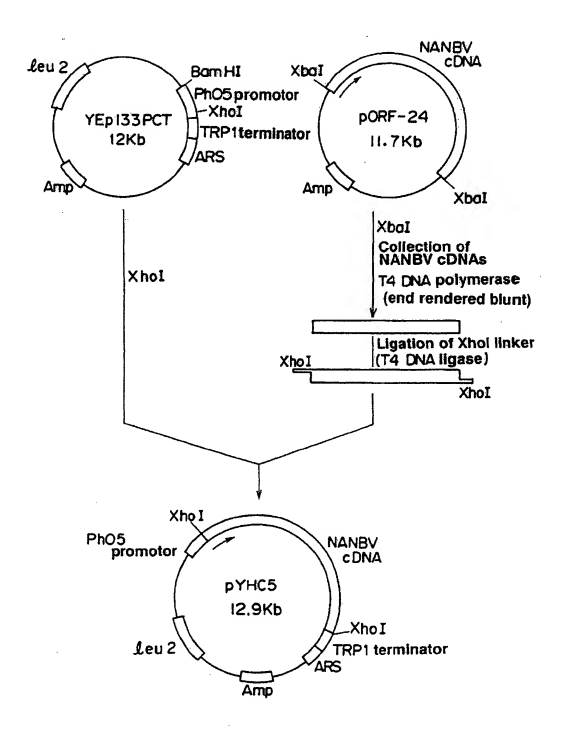


FIG. 6

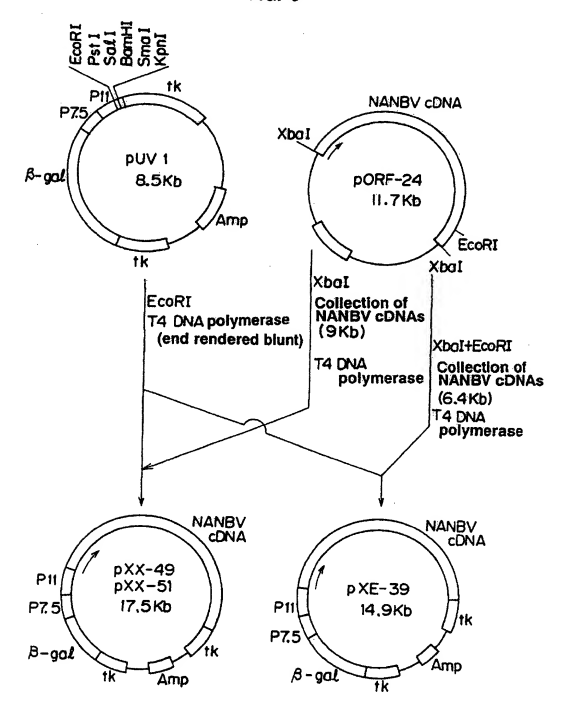


FIG. 7

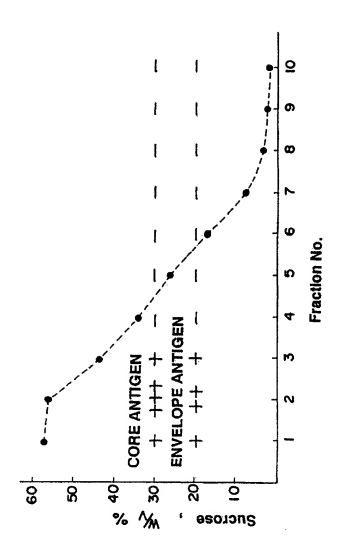
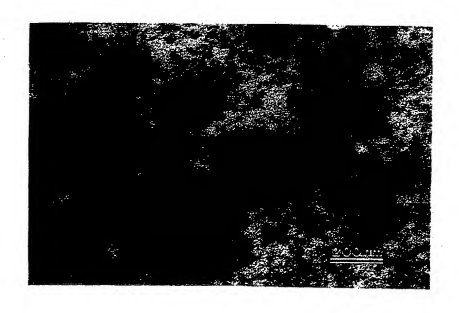


FIG. 8



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